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(54) Title: OUTER MEMBRANE PROTEIN OF EHRLICHIA CANIS AND EHRLICHIA CHAFFEENSIS

(57) Abstract

The present invention relates to diagnostic tools for veterinary and human use which are used for serodiagnosing ehrlichiosis in mammals, particularly in members of the Canidae family and in humans. The present invention also provides polynucleotides which encode the outer membrane proteins of E. chafeensis. The polynucleotides encode an OMP-1 family of proteins of E. chafeensis and P30 family of proteins of E. canis. The present invention also provides the following isolated proteins of E. chafeensis OMP-1, OMP-1A, OMP-1B, OMP-1C, OMP-1D, OMP-1B, OMP-1B, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, and OMP-1Z, referred to hereinafter collectively as the "OMP family". The present invention also provides the following isolated proteins of E. canis P30, P30-a, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10, referred to hereinafter as the P30 family. The present invention also relates to an assay for diagnosing ehrlichiosis in humans using a recombinant outer membrane protein of E. chafeensis, particularly OMP-1. The present invention also relates to an assay for diagnosing ehrlichiosis in humans and members of the family Canidae using a recombinant outer membrane protein of E. canis, particularly P30.

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OUTER MEMBRANE PROTEIN OF EHRLICHIA CANIS AND EHRLICHIA CHAFFEENIS

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BACKGROUND OF THE INVENTION

The ehrlichiae are obligate intracellular bacteria that infect circulating leucocytes. Ehrlichia chafeensis infects the monocytes and macrophages in humans and causes human monocytic ehrlichiosis. The clinical manifestations of ehrlichiosis in humans are nonspecific and similar to Rocky Mountain spotted fever. The clinical manifestations include fever, chills, headache mylagia or vomiting and weight loss. Most patients have a history of tick exposure.

Ehrlichia canis infects and causes ehrlichiosis in animals belonging to the family Canidae. Canine ehrlichiosis consists of an acute and a chronic phase. The acute phase is characterized by fever, serous nasal and ocular discharges, anorexia, depression, and loss of weight. The chronic phase is characterized by severe pancytopenia, epistaxis, hematuria, blood in feces in addition to more severe clinical signs of the acute disease. If treated early during the course of the disease, dogs respond well to doxycycline. However, chronically infected dogs do not respond well to the antibiotic. Therefore, early diagnosis is very important for treating canine ehrlichiosis.

The primary diagnostic test for diagnosing canine ehrlichiosis and human ehrlichiosis is the indirect fluorescent antibody (IFA) test. This test uses the etiologic agent Ehrlichia canis to diagnose canine ehrlichiosis. The IFA test uses Ehrlichia chafeensis as antigen for diagnosing human ehrlichiosis. The IFA test has, however, serious limitations. The IFA test is subject to false positives because the antigens are made of whole infected cells which comprise many nonspecific proteins which will cross-react with sera from some patients. The IFA test is also subject to false negatives because IFA antigens are unstable and may become inactivated during storage. In addition the IFA test requires a special equipment to perform the test. For example, the IFA test requires a tissue culture system for growing the bacterium that are used to prepare the antigen slides, a fluorescent microscope, and trained persons to evaluate the serum reactivity to the bacterial antigen on the slide.

Tools which permit simpler, more rapid, and objective serodiagnosis of canine ehrlichiosis or human ehrlichiosis are desirable.

SUMMARY OF THE INVENTION

The present invention relates to improved diagnostic tools for veterinary and human use which are used for serodiagnosing ehrlichiosis in mammals, particularly in members of the Canidae family and in humans.

The present invention also provides polynucleotides or nucleic acids which encode the outer membrane proteins of E. chafeensis. The OMP-1 polynucleotide encodes an OMP-1 protein of E. chafeensis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG.3B, SEQ ID NO: __. The OMP-1B polynucleotide encodes an OMP-1B protein of E.

chafeensis having a molecular weight of about 28.2 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 4B, SEQ ID NO: __. The OMP-1C polynucleotide encodes an OMP-1C protein of E. chafeensis having a molecular weight of about 27.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 5B, SEQ ID NO: __. The OMP-1D polynucleotide encodes an OMP-1D protein of E. chafeensis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 6B, SEQ ID NO: __. The OMP-1E polynucleotide encodes an OMP-1E protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 7B, SEQ ID NO: __. The OMP-1F polynucleotide encodes an OMP-1F protein of E. chafeensis having a molecular weight of about 27.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 8B, SEQ ID NO: __. The OMP-1A polynucleotide encodes an OMP-1A protein of E. chafeensis having a molecular weight of about 29.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 9B, SEQ ID NO: __. The OMP-1R polynucleotide encodes an OMP-1R protein of E. chafeensis having a molecular weight of at least 23 kDa and comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 10B, SEQ ID NO: __. The OMP-1S polynucleotide encodes an OMP-1S protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 11B, SEQ ID NO: __. The OMP-1T polynucleotide encodes an OMP-1T protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 12B, SEQ ID NO: __. The OMP-1U polynucleotide encodes an OMP-1U protein of E. chafeensis having a molecular weight of about 30.6 kDa and an amino acid sequence which is at least 85% homologous to amino acid sequence shown in FIG. 13B, SEQ ID NO: __. The OMP-1V polynucleotide encodes an OMP-1V protein of E. chafeensis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 14B, SEQ ID NO: __. The OMP-1W polynucleotide encodes an OMP-1W protein of E. chafeensis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 15B, SEQ ID NO: __. The OMP-1X polynucleotide encodes an OMP-1S protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 16B, SEQ ID NO: __. The OMP-1Y polynucleotide encodes an OMP-1Y protein of E. chafeensis having a molecular weight about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 17B, SEQ ID NO: __. The OMP-1Z polynucleotide encodes an OMP-1Z protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 18B, SEQ ID NO: __.

The outer membrane proteins from E. chaffeensis, particularly a recombinant form of OMP-1, are immunogenic and, thus are useful for preparing antibodies. Such antibodies are useful for immunolabeling isolates of E. chafeensis and for detecting the presence of E. chafeensis in body fluids, tissues, and particularly in monocytes and macrophages. The isolated outer membrane proteins, particularly OMP-1, are also useful for

detecting antibodies to E. chafeensis in the blood of patients with clinical signs of ehrlichiosis. The isolated outer membrane protein, particularly OMP-1, are also useful immunogens for raising antibodies that are capable of reducing the level of infection in an immunized mammal that has been infected with E. chafeensis. The isolated membrane proteins are also useful in a vaccine for protecting against infection with E. chafeensis.

The present invention also relates to isolated polynucleotides which encode 30 kDa outer membrane proteins from Ehrlichia canis. The proteins are designated P30 and P30a. The proteins, particularly P30, are immunogenic and are, thus, useful for preparing antibodies that are useful for immunolabeling isolates of E. canis. The P30 protein is also useful for diagnosing canine ehrlichiosis in mammals, particularly in members of the family Canidae, most particularly in dogs and for diagnosing infections with E. chafeensis in humans. The P30 protein is also a useful immunogen for raising antibodies that reduce the level of infection in an immunized mammal that has been infected with E. canis. The P30 protein is also useful in a vaccine for protecting animals against infection with E. canis.

The present invention also provides the following isolated proteins of E. chafeensis OMP-1 (also known as p28), OMP-1A, OMP-1B, OMP-1C, OMP-1D, OMP-1E, OMP-1F, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, and OMP-1Z, referred to hereinafter collectively as the "OMP family". The present invention also provides the following isolated proteins of E. canis P30, P30-a, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10, referred to hereinafter as the P30 family.

The present invention also relates to an assay for diagnosing ehrlichiosis in humans using a recombinant outer membrane protein of E. chafeensis, particularly OMP-1. The present invention also relates to an assay for diagnosing ehrlichiosis in humans and members of the family Canidae using a recombinant outer membrane protein of E. canis, particularly P30.

Brief Description of the Figures

- FIG. 1. shows the DNA sequence of and the amino acid sequence encoded by the E. chafeensis (p28) gene cloned in pCRIIp28. The N-terminal amino acid sequence of native omp-1 protein (P28) determined chemically is underlined. Five amino acid residues at the N terminus of P28 which were not included in the p28 gene, are indicated by boldface. Arrows indicate annealing positions of the primer pair designed for PCR
- FIG. 2. shows the restriction map of 6.3-kb genomic DNA including the *omp-1* gene copies in *E. chafeensis*. The four DNA fragments were cloned from the genomic DNA (pPS2.6, pPS3.6, pEC2.6, and pEC3.6). A recombinant plasmid pPS2.6 has an overlapping sequence with that of pEC3.6. The closed boxes at the bottom show PCR-amplified fragments from the genomic DNA for confirmation of the overlapping area. Open boxes at the top indicate open reading frames (ORF) of *omp-1* gene copies with direction by arrows. Open boxes at the bottom show DNA fragments subcloned for DNA sequencing.
- FIG. 3B shows one embodiment of the OMP-1 protein; FIG. 3A shows one embodiment of the OMP-1 polynucleotide.
- FIG. 4B shows one embodiment of the OMP-1B protein, FIG. 4A shows one embodiment of the OMP-1B polynucleotide

FIG. 5A shows one embodiment of the OMP-1C polynucleotide; FIG 5B shows one embodiment of the OMP-1C protein.

- FIG. 6B shows one embodiment of the OMP-1D protein; FIG. 6A shows one embodiment of the OMP-1D polynucleotide.
- FIG. 7A shows one embodiment of the OMP-1E protein; FIG 7B shows one embodiment of the OMP-1E polynucleotide.
- FIG. 8A shows one embodiment of the OMP-1F protein; FIG 8 B shows one embodiment of the OMP-1F polynucleotide.
- FIG. 9B shows one embodiment of the OMP-1A protein, FIG 9A shows one embodiment of the OMP-1A polynucleotide.
- FIG. 10 B shows one embodiment of a portion of the OMP-1R protein, FIG 10A shows one embodiment of an OMP-1R polynucleotide encoding such polypeptide.
- FIG. 11 B shows one embodiment of a portion of the OMP-1S protein, FIG 11A shows one embodiment of the OMP-1S polynucleotide encoding such polypeptide.
- FIG. 12 B shows one embodiment of a portion of the OMP-1T protein, FIG 12A shows one embodiment of the OMP-1T polynucleotide encoding such polypeptide.
- FIG. 13 B shows one embodiment of the OMP-1U protein, FIG 13A shows one embodiment of the OMP-1U polynucleotide.
- FIG. 14 B shows one embodiment of the OMP-1V protein, FIG 14A shows one embodiment of the OMP-1V polynucleotide.
- FIG. 15 B shows one embodiment of the OMP-1W protein, FIG 15A shows one embodiment of the OMP-1W polynucleotide.
- FIG. 16 B shows one embodiment of the OMP-1X protein, FIG 16A shows one embodiment of the OMP-1W polynucleotide.
- FIG. 17 B shows one embodiment of the OMP-1Y protein, FIG 17A shows one embodiment of the OMP-1Y polynucleotide.
- FIG. 18 B shows one embodiment of the OMP-1Z protein, FIG 18A shows one embodiment of the OMP-1Z polynucleotide.
- FIG. 19 B shows one embodiment of the P30 protein, FIG 19A shows one embodiment of the P30 polynucleotide.
- FIG. 20 B shows one embodiment of the P30a protein, FIG 20A shows one embodiment of the p30A polynucleotide.
- FIG. 21 B shows one embodiment of the P30-1 protein, FIG 21A shows one embodiment of the p30-1 polynucleotide.
- FIG. 22 B shows one embodiment of the P30-2 protein, FIG 22 A shows one embodiment of the p30-2 polynucleotide.

FIG. 23 B shows one embodiment of the P30-3 protein, FIG 23 A shows one embodiment of the p30-3 polynucleotide.

- FIG. 24 B shows one embodiment of the P30-4 protein, FIG 22 A shows one embodiment of the p30-4 polynucleotide.
- FIG. 25 B shows one embodiment of the P30-5 protein, FIG 22 A shows one embodiment of the p30-5 polynucleotide.
- FIG. 26 B shows one embodiment of the P30-6 protein, FIG 26 A shows one embodiment of the p30-6 polynucleotide.
- FIG. 27 B shows one embodiment of the P30-7 protein, FIG 27 A shows one embodiment of the p30-7 polynucleotide.
- FIG. 28 B shows one embodiment of the P30-8 protein, FIG 28 A shows one embodiment of the p30-8 polynucleotide.
- FIG. 29 B shows one embodiment of a portion of the P30-9 protein, FIG 29 A shows one embodiment of the p30-9 polynucleotide encoding such polypeptide.
- FIG. 30 B shows one embodiment of a portion of the P30-10 protein, FIG 30 A shows one embodiment of the p30-10 polynucleotide encoding such polypeptide.
- FIG. 31 depicts the amino acid sequences alignment of seven E. chafeensis OMP-1s and Cowdria ruminantium MAP-1. Aligned positions of identical amino acids with OMP-IF are shown with dots. The sequence of C. ruminantium MAP-1 is from the report of Van Vliet et al (1994) Molecular cloning, sequence analysis, and expression of the gene encoding the immunodominant 32-kilodalton protein of Cowdria ruminantium. Infect. Immun. 62:1451-1456. Gaps indicated by dashes were introduced for optimal alignment of all proteins. Bars indicates semivariable region (SV) and three hypervariable regions (HV1, HV2, and HV3).

DETAILED DESCRIPTION OF THE INVENTION

Isolated Polynucleotides Encoding OMP-1.OMP-1A. OMP-1B. OMP-1C. OMP-1D, OMP-1F and the OMP from E. Canis

In one aspect, the present invention, provides isolated polynucleotides that encode the outer membrane proteins, OMP-1 (or p28), OMP-1B, OMP-1C, OMP-1D, OMP-1E, OMP-1F, OMP-1A, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, OMP-1Y and OMP-1Z from E. chafeensis and the outer membrane proteins P30, P30-a, P-30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10 from E. Canis or an immunogenic fragment thereof.

The polynucleotide is single stranded or double stranded. The polynucleotide may be a DNA or RNA molecule, preferably a DNA molecule, and comprises a sequence which codes for the respective outer membrane protein. Preferably, the polynucleotide encodes at least the mature form of outer membrane protein. The polynucleotide optionally further comprises a leader sequence and encode an outer membrane preprotein that is

processed in the cell to form the mature protein. The polynucleotide of the present invention may also be fused in frame to a marker sequence which allows for purification of the corresponding outer membrane protein.

The OMP-1 polynucleotide encodes an OMP-1 protein of E. chafeensis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 3B SEQ ID NO: __; Figure 3B shows one embodiment of the OMP-1 protein, Figure 3A shows one embodiment of the OMP-1 polynucleotide. The OMP-1B polynucleotide encodes an OMP-1B protein of E. chafeensis having a molecular weight of about 28.2 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 4B SEQ ID NO: __; Figure 4B shows one embodiment of the OMP-1B protein, Figure 4A shows one embodiment of the OMP-1B polynucleotide. The OMP-1C polynucleotide encodes an OMP-1C protein of E. chafeensis having a molecular weight of about 27.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 5B SEQ ID NO: __; Figure 5B shows one embodiment of the OMP-1C protein, Figure 5A shows one embodiment of the OMP-1C polynucleotide. The OMP-1D polynucleotide encodes an OMP-1D protein of E. chafeensis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 6B SEQ ID NO: __; Figure 6B shows one embodiment of the OMP-1D protein, Figure 6A shows one embodiment of the OMP-1D polynucleotide. The OMP-1E polynucleotide encodes an OMP-1E protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 7B SEQ ID NO: __; Figure 7B shows one embodiment of the OMP-1E protein, Figure 7A shows one embodiment of the OMP-1E polynucleotide. The OMP-1F polynucleotide encodes an OMP-1F protein of E. chafeensis having a molecular weight of about 27.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 8B SEQ ID NO: ___; Figure 8B shows one embodiment of the OMP-1F protein, Figure 8A shows one embodiment of the OMP-1F polynucleotide. The OMP-1A polynucleotide encodes an OMP-1A protein of E. chafeensis having a molecular weight of about 29.6 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 9B SEQ ID NO: __; Figure 9B shows one embodiment of the OMP-1A protein, Figure 9A shows one embodiment of the OMP-1A polynucleotide. The OMP-1R polynucleotide encodes an OMP-1R protein of E. chafeensis having a molecular weight of at least 23 kDa and comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 10B SEQ ID NO: __; Figure 10B shows one embodiment of a portion of the OMP-1R protein, Figure 10A shows one embodiment of the OMP-1R polynucleotide encoding such polynucleotide. The OMP-1S polynucleotide encodes an OMP-1S protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 11B SEQ ID NO: __; Figure 11B shows one embodiment of a portion of the OMP-1S protein, Figure 11A shows one embodiment of the OMP-1S polynucleotide encoding such polypeptice. The OMP-1T polynucleotide encodes an OMP-1T protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG.12B SEQ ID NO: __; Figure 12B shows one embodiment of a portion of the OMP-1T protein, Figure 12B shows one embodiment of a polynucleotide encoding such polypeptide. The OMP-1U polynucleotide encodes an

OMP-1U protein of E. chafeensis having a molecular weight of about 30.6 kDa and an amino acid sequence which is at least 85% homologous to amino acid sequence shown in FIG. 13B SEQ ID NO: __; Figure 13B shows one embodiment of the OMP-1U protein, Figure 13A shows one embodiment of the OMP-1U polynucleotide. The OMP-1V polynucleotide encodes an OMP-1V protein of E. chafeensis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 14B SEQ ID NO: ___; Figure 14B shows one embodiment of the OMP-1V protein, Figure 14A shows one embodiment of the OMP-1V polynucleotide. The OMP-1W polynucleotide encodes an OMP-1W protein of E. chafeensis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 15B SEQ ID NO: __; Figure 15B shows one embodiment of the OMP-1W protein, Figure 15A shows one embodiment of the OMP-1W polynucleotide. The OMP-1X polynucleotide encodes an OMP-1S protein of E. chafeensis having a molecular weight of about 27.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 16B SEQ ID NO: __; Figure 16B shows one embodiment of the OMP-1X protein, Figure 16A shows one embodiment of the OMP-1X polynucleotide. The OMP-1Y polynucleotide encodes an OMP-1Y protein of E. chafeensis having a molecular weight about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 17B SEQ ID NO: __; Figure 17B shows one embodiment of the OMP-1Y protein, Figure 17A shows one embodiment of the OMP-1Y polynucleotide. The OMP-1Z polynucleotide encodes an OMP-1Z protein of E. chafeensis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 18B SEQ ID NO: Figure 18B shows one embodiment of a portion of the OMP-1Z protein, Figure 18A shows one embodiment of an OMP-1Z polynucleotide encoding such polypeptide.

The p30 polynucleotide encodes a P30 protein of E. canis having a molecular weight of about 28.8 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 19B SEQ ID NO: ___; Figure 19B shows one embodiment of the P30 protein, Figure 19A shows one embodiment of the p30 polynucleotide. The p30A polynucleotide encodes a P30a protein of E. canis having a molecular weight of about 29.1 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 20B SEQ ID NO: ___; Figure 20B shows one embodiment of the P30a protein, Figure 20A shows one embodiment of the p30A polynucleotide. The p30-1 polynucleotide encodes a P30-1 protein of E. canis having a molecular weight of about 27.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 21B SEQ ID NO: __; Figure 21B shows one embodiment of the P30-1 protein, Figure 21A shows one embodiment of the p30-1 polynucleotide. The p30-2 polynucleotide encodes a P30-2 protein of E. canis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 22B SEQ ID NO: __; Figure 22B shows one embodiment of the P30-2 protein, Figure 22A shows one embodiment of the p30-2 polynucleotide. The p30-3 polynucleotide encodes a P30-3 protein of E. canis having a molecular weight of about 28.7 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 23B SEQ ID NO: ___, Figure 23B shows one embodiment of the P30-3 protein, Figure 23A shows one embodiment of the p30-3 polynucleotide. The p30-4 polynucleotide

encodes a P30-4 protein of E. canis having a molecular weight of about 28.0 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 24B SEQ ID NO: __; Figure 24B shows one embodiment of the P30-4 protein, Figure 24A shows one embodiment of the p30-4 polynucleotide. The p30-5 polynucleotide encodes a P30-5 protein of E. canis having a molecular weight of about 29.4 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 25B SEQ ID NO: __; Figure 25B shows one embodiment of the P30-5a protein, Figure 25A shows one embodiment of the p30-5a polynucleotide. The p30-6 polynucleotide encodes a P30-6 protein of E. canis having a molecular weight of about 29.5 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 26B SEQ ID NO: ___; Figure 26B shows one embodiment of the P30-6 protein, Figure 26A shows one embodiment of the p30-6 polynucleotide. The p30-7 polynucleotide encodes a P30-7 protein of E. canis having a molecular weight of about 29.9 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 29B SEQ ID NO: __; Figure 29B shows one embodiment of the P30-7 protein, Figure 29A shows one embodiment of the p30-7 polynucleotide. The p30-8 polynucleotide encodes a P30-8 protein of E. canis having a molecular weight of about 30.3 kDa and an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 28B SEQ ID NO: __; Figure 28B shows one embodiment of the P30-8 protein, Figure 28A shows one embodiment of the p30-8 polynucleotide. The p30-9 polynucleotide encodes a P30-9 protein of E. canis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 29B SEQ ID NO: ___; Figure 29B shows one embodiment of a portion of the P30-9 protein, Figure 29A shows one embodiment of the p30-9 polynucleotide encoding such polypeptide. The p30-10 polynucleotide encodes a P30-10 protein of E. canis comprising an amino acid sequence which is at least 85% homologous to the amino acid sequence shown in FIG. 30B SEQ ID NO: __; Figure 30B shows one embodiment of a portion of the P30-10 protein, Figure 30A shows one embodiment of the p30-10 polynucleotide encoding such polypeptide.

The polynucleotides encoding an E. chafeensis outer membrane protein or an E. canis outer membrane protein have a sequence that is at least 85%, preferably at least 90%, more preferably at least 95% homologous to or similar to the amino acid sequences shown in Figures 3B through 30B, and thus embrace polynucleotides encoding outer membrane proteins from different strains of E. chafeensis and E. canis. The polynucleotides encode an outer membrane protein whose conserved regions collectively are at least 90%, preferably at 95%, more preferably at least 97% homologous to the conserved regions of the amino acid sequences of the present invention. The outer membrane proteins of E. chafeensis and E. canis have six conserved regions, which are separated by one semivariable region and three hypervariable regions. The conserved regions of the outer membrane proteins OMP-1, OMP-1B, OMP1-C, OMP-1D, OMP1-F are depicted in Fig. 31. Preferably, the amino acid sequence of the outer membrane proteins of E. chafeensis and E. canis are at least 30% divergent from the amino acid sequence of MAP-1. Such sequences include allelic, strain variants and other amino acid sequence variants (e.g., including "muteins" or "mutant proteins"), whether naturally-occurring or biosynthetically produced. As used herein, "amino acid sequence homology" is understood to mean amino acids are conserved amino acids as defined by sequences share identical or similar amino acids, where similar amino acids are conserved amino acids as defined by

Dayoff et al., Atlas of Protein Sequence and Structure; vol. 5, Supp. 3, pp. 345-362 (M. O. Dayoff, ed., Nat'l BioMed. Research Fdn., Washington D.C. 1978.) Thus, a candidate sequence sharing 85% amino acid sequence homology with a reference sequence requires that, following alignment of the candidate sequence with the reference sequence, 85% of the amino acids in the candidate sequence are identical to the corresponding amino acid in the reference sequence, or constitute a conserved amino acid change thereto. "Amino acid sequence identity" is understood to require identical amino acids between two aligned sequences. Thus, a candidate sequence sharing 85% amino acid identity with a reference sequence requires that, following alignment of the candidate sequence with the reference sequence, 85% of the amino acids in the candidate sequence are identical to the corresponding amino acid in the reference sequence.

As used herein, all homologies and identities are calculated using the amino acid sequences shown in the cited Figure or SEQ ID NO as the reference sequence. Thus, to determine whether an amino acid sequence is 85% homologous to OMP-1, one uses the amino acid sequence shown in Fig. ___, SEQ ID NO: ___ as a reference.

Also as used herein, sequences are aligned for homology and identity calculations using the method of the software basic local alignment search tool in the BLAST network service (the National Center for Biotechnology Information, Bethesda, MD) which employs the method of Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. (1990) J. Mol. Biol. 215, 403-410. Identities are calculated by the Align program (DNAstar, Inc.) In all cases, internal gaps and amino acid insertions in the candidate sequence as aligned are ignored when making the homology/identity calculation.

In another aspect, the present invention provides a nucleotide sequence encoding a polypeptide which comprises a fragment of the OMP1 protein, hereinafter referred to as "rP28". The rP28 polypeptide weighs approximately 31 kDa and comprises all but of the first 5 amino acids of mature OMP-1 protein. The rP28 polypeptide comprises the amino acid sequence extending from amino acid 6 through amino acid 251 of the amino acid sequence shown in Fig.1, SEQ ID NO. The present invention also embraces polypeptides where one or more of the amino acids in the sequence extending from amino acid 1 or 6 through amino acid 251 Fig. 1 are replaced by conservative amino acid residues. The present invention also relates to derivatives of rP28 that have an amino acid sequence identity of at least 85%, more preferably at least 90%, and most preferably of at least 95% with the amino acid sequence extending from amino acid 1 or 6 through amino acid 251 of the protein and which derivative binds to antibodies in sera from humans infected with E. chafeensis.

The polynucleotides are useful for producing the outer membrane proteins of E. chafeensis and E. canis. For example, an RNA molecule encoding the outer membrane protein OMP-1 is used in a cell-free translation systems to prepare OMP-1. Alternatively, a DNA molecule encoding the outer membrane protein is introduced into an expression vector and used to transform cells. Suitable expression vectors include for example chromosomal, nonchromosomal and synthetic DNA sequences, e.g., derivatives of SV40, bacterial plasmids, phage DNAs; yeast plasmids, vectors derived from combinations of plasmids and phage DNAs, viral DNA such as vaccinia, adenovirus, fowl pox virus, and pseudorabies. The DNA sequence is introduced into the expression vector by conventional procedures.

Accordingly, the present invention also relates to recombinant constructs comprising one or more of the polynucleotide sequences. Suitable constructs include, for example, vectors, such as a plasmid, phagemid, or viral vector, into which a sequence that encodes the outer membrane protein has been inserted. In the expression vector, the DNA sequence which encodes the outer membrane protein is operatively linked to an expression control sequence, i.e., a promoter, which directs mRNA synthesis. Representative examples of such promoters, include the LTR or SV40 promoter, the E.coli lac or trp, the phage lambda PL promoter and other promoters known to control expression of genes in prokaryotic or eukaryotic cells or in viruses. The promoter may also be the natural promoter of the outer membrane protein coding sequence. The expression vector also contains a ribosome binding site for translation initiation and a transcription terminator. Preferably, the recombinant expression vectors also include an origin of replication and a selectable marker, such as for example, the ampicillin resistance gene of E. coli to permit selection of transformed cells, i.e. cells that are expressing the heterologous DNA sequences. The polynucleotide sequence encoding the outer membrane protein is incorporated into the vector in frame with translation initiation and termination sequences. Optionally, the sequence encodes a fusion outer membrane protein which includes an N-terminal or C-terminal peptide or tag that stabilizes or simplifies purification of the expressed recombinant product. Representative examples of such tags include sequences which encode a series of histidine residues, the Herpes simplex glycoprotein D, or glutathione S-transferase.

Polynucleotides which encode portions of the outer membrane proteins of E. chafeensis and E. canus are useful as probes for isolating and identifying E. chafeensis genes and E. canis genes, particularly full-length genes from new strains or isolates of E. chafeensis and E. canis.

The Outer Membrane Proteins of E. chafeensis and E. Canis

Preparing the Outer Membrane Proteins

4

In addition to the outer membrane proteins OMP-1, OMP-1B, OMP-1C, OMP-1D, OMP-1 E, and OMP-1F, OMP-1R, OMP-1S, OMP-1T, OMP-1U, OMP-1V, OMP-1W, OMP-1X, OMP-1Y, and OMP-1Z from E. chafeensis and the proteins P30, P30A, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6, P30-7, P30-8, P30-9, and P30-10 from E. Canis, the present inventions embraces non-naturally occurring allelelic forms or derivatives of the outer membrane proteins; where one or more of the amino acids have been replaced by conservative amino acid residues, typically by using direct synthesis or recombinant techniques.

The outer membrane proteins of the present invention are synthetically produced by conventional peptide synthesizers. The outer membrane proteins are also produced using cell-free translation systems and RNA molecules derived from DNA constructs that encode the outer membrane protein. Alternatively, the outer membrane protein is made by transfecting host cells with expression vectors that comprise a DNA sequence which encodes the outer membrane protein and then inducing expression of the outer membrane protein in the host cells.

The outer membrane protein is expressed in suitable host cells, preferably bacteria, under the control of suitable promoters. Host cells are transformed with the expression vectors of this invention and cultured in conventional nutrient media. Such media optionally contains additional compounds, such as for example

compounds that induce promoters, such as for example isopropyl-β-D-thiogalactoside which induces the Lac promoter, or compounds, such as for example, ampicillin, which allows for selection of transformants.

Following transformation of the suitable host strain and growth of the host strain to an appropriate cell density, the cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained for further purification of the outer membrane protein. Such purification usually involves salting-out of the protein fraction, and one or more chromatography steps, including aqueous ion exchange chromatography, size exclusion chromatography steps, and high performance liquid chromatography (HPLC). Preparation of Antibodies

The isolated outer membrane proteins, particularly the recombinant forms of the outer membrane proteins, are used as immunogens to produce antibodies immunospecific for the corresponding protein. The term "immunospecific" means the antibodies have substantially greater affinity for the protein used as an immunogen than for other proteins. Such antibodies are generated using conventional techniques by administering the respective outer membrane protein or a portion thereof, i.e., the recombinant polypeptide, to an animal, preferably a nonhuman. collecting blood from the immunized animals and isolating the serum and/or the IgG fraction from the blood. Monoclonal antibodies are prepared by injecting animals with the immunogens, extracting antibody-producing B cells from the animal, fusing the B cells with a myeloma cells to produce hybridomas, obtaining the monoclonal antibodies from the hybridomas.

Antibodies to the outer membrane proteins of E. chafeensis and E. canis are useful research tools for identifying cells, particularly monocytes, infected with E.chafeensis or E. canis and for purifying the corresponding outer membrane protein of E.chafeensis or E. Canis from partially purified preparations by affinity chromatography. Such antibodies are also useful for identifying bacterial colonies, particularly colonies of genetically-engineered bacteria, that are expressing the major outer membrane protein.

Diagnostic Method

The present invention also provides a method for detecting antibodies to the E. chafeensis or E. canis in a sample of a bodily fluid from a patient. The method comprises providing an isolated outer membrane protein of E. chafeensis or E. canis, particularly a recombinant form of the isolated protein, contacting the outer membrane protein or polypeptide with a sample taken from the patient; and assaying for the formation of a complex between the outer membrane protein or polypeptide and antibodies in the sample. For ease of detection, it is preferred that the isolated protein or polypeptide be attached to a substrate such as a column, plastic dish, matrix, or membrane, preferably nitrocellulose. The sample may be a tissue or a biological fluid, including urine, whole blood, or exudate, preferably serum. The sample may be untreated, subjected to precipitation, fractionation, separation, or purification before combining with the isolated protein or peptide. Interactions between antibodies in the sample and the isolated protein or peptide are detected by radiometric, colorimetric, or fluorometric means, size-separation, or precipitation. Preferably, detection of the antibody-outer membrane protein complex is by addition of a secondary antibody that is coupled to a detectable tag, such as for example, an enzyme, fluorophore, or chromophare. Formation of the complex is indicative of the presence of anti-E chafeensis or anti-E canis antibodies,

either IgM or IgG, in the patient. Thus, the method is used to determine whether a patient is infected with E. chafeensis or E. canis.

Preferably, the method employs an enzyme-linked immunosorbent assay (ELISA) or a Western immunoblot procedure. Such methods are relatively simple to perform and do not require special equipment as long as membrane strips are coated with a high quality antigen. Accordingly, it is more advantageous to use a recombinant form of the outer membrane protein of E. chafeensis or E. canis since such proteins, typically, are more pure and consistent in quality than a purified form of such protein.

Immunogenic Composition

The present invention also relates to immunogenic compositions comprising one or more of the isolated outer membrane proteins of E. chafeensis and a pharmaceutically acceptable adjuvant and to immunogenic compositions comprising an isolated P30 protein of E. canis and a pharmaceutically acceptable adjuvant, which, preferably, enhances the immunogenic activity of the outer membrane protein in the host animal.

Preparation of a Polynucleotide which Encodes OMP-1(P28)

A. Isolation of the Outer Membrane Proteins

E. chafeensis Arkansas strain and E. canis Oklahoma strain were cultivated in the DH82 dog macrophage cell line and purified by Percoll density gradient centrifugation. Purified ehrlichiae (100 μg) were suspended with 10 mM sodium phosphate buffer, pH 7.4, containing 0.1% Sodium N-lauroyl sarcosine (Sarkosyl) [Sigma, St. Louis, MO], 50 μg/ml each Dnase I (Sigma) and Rnase A (Sigma), and 2.5 mM MgCl₂. After incubation at 37° for 30 min, the sample was separated by centrifugation at 10,000 x g for 1 h into the soluble supernatant and the insoluble precipitate. The insoluble pellet was resuspended 2 to 3 times with 0.1% Sarkosyl and centrifuged. The final pellet was analyzed by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) and by electron microscopy.

Transmission electron microscopy revealed that the purified ehrlichial fraction consists of a mixture of electron dense and light forms of *E. chafeensis* with slight disintegration of inner membrane. Ehrlichiae were not surrounded with the host inclusion membrane. Various sizes of membrane vesicles (< 1 µm) without significant ribosomes or nuclear materials were observed in the Sarkosyl-insoluble fraction from the organism. Succinic dehydrogenase (inner membrane marker enzyme of gram negative bacteria) activities were at less than the detection limit (1 n moles / min / mg of protein) in the Sarkosyl-insoluble fraction compared to approximately 10 n moles / min / mg of protein in the Percoll-purified organisms, suggesting that the insoluble fraction primarily consisted of the outer membrane of *E. chafeensis*.

Analysis of the Sarkosyl-soluble, and insoluble fraction of *E. chafeensis* by SDS-PAGE suggested that proteins of 30-kDa range in the insoluble fraction represent the major outer membrane proteins of this organism. Analysis of the Sarkosyl-soluble, and insoluble fraction of *E. canis* by SDS-PAGE suggested that proteins of 30-kDa range in the insoluble fraction represent the major outer membrane proteins of this organism also. *E. canis* was

antigenically cross reactive with *E. chafeensis*. These findings indicate that the 30-kDa range proteins represent the major outer membrane proteins of these two *Ehrlichia* spp.

To improve resolution of the outer membrane proteins, proteins in the Sarkosyl-insoluble pellet prepared from 400 µg of purified *E. chafeensis* were separated by a reversed-discontinuous (Rd) SDS-PAGE (2.5-cm-long 17% gel on top of 11-cm-long 12% gel). At least five proteins of 30-kDa range in *E. chafeensis* (P23, P25, P27, P28, and P29) were resolved from the Sarkosyl-insoluble proteins.

B. Cloning and sequencing of the p28 gene

The	portion of the	nembrane contair	ning bound prot	eins was exc	ised and analyzed	with an	Applied
Biosystems p	rotein sequencer	(Model 470). The	N-terminal ami	no acid seque	nce of P28 was det	termined a	is DPA
GSGING	NFYSGKY	M P, SEQ IN N	O Ba	sed on 6th to	12th amino acids of	of this seq	uence, a
forward	primer,	FECH1,	having	the	sequence:		5'-
CGGGATCC	GAATTCGG(A	T/G/C)AT(A/T/C)AA(T/C)GG(A	T/G/C)AA(T/	C)TT(T/C)TA-3'.	SEQ 1	ID NO
was	designed. Ami	no acids at the I	to 5 positions o	f the N termi	nus of P28 were no	ot include	d in this
primer design	n. For insertion	into an expression	vector, a 14-bp	sequence (ur	iderlined) was adde	ed at the S	5' end of
primer to cre	ate an EcoRI and	i a BamHi site. Th	ne reverse prime	r, RECH2, w	hich includes a Not	I site at th	ne 5' end
for ligation i	nto an expression	n vector had the	sequence: 5'-A	.GCGGCCGC	TTA(A/G)AA(T/C)A(C/G) ((A/G)AA
		EQ ID NO					

Genomic DNA of *E. chafeensis* was isolated from purified organisms. PCR amplification with FECH1 and RECH2 primers was performed using a Perkin-Elmer Cetus DNA Thermal Cycler (model 480). A 0.8-kb amplified product was cloned in the pCRII vector of a TA closing kit, as described by the manufacturer (Invitrogen Co., San Diego, CA). The clone obtained was designated pCRII*p28*. Both strands of the inserted DNA were sequenced by a dideoxy-termination method with an Applied Biosystems 373A DNA sequencer.

The 0.8-kb DNA fragment, cloned in pCRIIp28, had an open reading frame (ORF) of 756 bp encoding a 251-amino acid recombinant protein (including both PCR primer regions) with a molecular mass of 27,685 Da. The nucleotide sequence of the open reading frame, SEQ ID NO: , and the amino acid sequence of the polypeptide of the OMP-1 protein, SEQ ID NO ___, are shown in Figs ____ and ____, respectively.

A DNA fragment comprising the p30 gene was prepared in a similar manner, i.e., by PCR amplification of genomic DNA of E. canis with the FECH1 and RECH2 primers.

Preparation of Polynucleotides which encode OMP 1A, OMP1B, OMP1-C, OMP-1D, OMP-1F, and OMP1-E

A. Southern blot analysis. Genomic DNA extracted from the purified *E. chafeensis* (200 ng each) was digested with restriction endonucleases, electrophoresed, and transferred to Hybond-N⁺ nylon membrane (Amersham, Arlington Heights, IL), by a standard method. The 0.8-kb p28 gene fragment from the clone pCRIIp28 was labeled with $[\alpha^{-12}P]$ dATP by the random primer method using a kit (Boehringer Mannheim, Indianapolis, IN) and the labeled fragment was used as a DNA probe. Hybridization was performed at 60°C in rapid hybridization buffer (Amersham) for 20 h. The nylon sheet was washed in 0.1 x SSC (1 x SSC containing 0.15M sodium chloride and

0.015M sodium citrate) with 1% SDS at 55°C and the hybridized probes were exposed to Hyperfilm (Amersham) at -80°C.

Genomic Southern blot analysis with several restriction enzymes resulted in one or more DNA fragment(s) of *E. chafeensis* which hybridized to ³²P-labeled *p28* gene probe. The restriction enzymes used did not cut within the *p28* gene portion of the pCRII*p28* insert. *Xba* I, *BgI* II, and *Kpn* I produced two bands, *Spe* I generated three bands, and *EcoR* V and *Pst* I produced multiple bands with different densities. *EcoR* I generated a broad band of 2.5 to 4kb. These *p28* homologous genes are designated as *omp-1* (outer membrane protein-1) family.

B. Cloning and sequencing of genomic copies of *E. chafeensis p28* gene. The *EcoR* I and *Pst* I fragments of DNA, detected by genomic Southern blot analysis as described above, were inserted into pBluescript II KS (+) vectors, and the recombinant plasmids were introduced into *E. coli* DH5α. Using the colony hybridization method with the ¹²P-labeled *p28* gene probe, four positive clones were isolated from the transformant. The positive clones were designated pEC2.6, pEC3.6, pPS2.6, and pPS3.6. These contained the ehrlichial DNA fragments of 2.6-kb (*EcoR* I), 3.6 kb (*EcoR* I), 2.6 kb (*Pst* I), and 3.6 kb (*Pst* I), respectively. The inserts of the clones pEC3.6 and pPS2.6 overlapped as shown in Fig._____. The overlapping area was further confirmed by PCR of *E. chafeensis* genomic DNA with two pairs of primer sets interposing the junctions of the four clones. The 1.1- to 1.6-kb DNA fragments of *HindIII-HindIII*, *HindIII-EcoRI*, or *XhoI-EcoRI* in the pEC2.6 and pEC3.6 were subcloned for sequencing. DNA sequencing was performed with suitable synthetic primers by dideoxy-termination method as described above.

Four DNA fragments from 2.6 to 3.6 kb were cloned from the *Eco*RI-digested and the *Pst*I-digested genomic DNA of *E. chafeensis* by colony hybridization with radiolabeled *p28* gene probe. The inserted DNA of the two recombinant clones, pEC3.6 and PPS2.6, were overlapped as shown in Fig. 7. Sequencing revealed one 5'-truncated ORF of 243 bp (designated *omp-1A*) and five complete ORF of 836-861 bp (designated *omp-1B* to *omp-1F*), which are tandemly-arrayed and are homologous to the *p28* gene (but are not identical), in the ehrlichial genomic DNA of 6,292 bp. The intergenic spaces were 581 bp between *omp-1A* and *omp-1B* and 260-308 bp among others. Putative promoter regions and ribosome-binding sites were identified in the noncoding regions.

Sequence analysis and GenBank accession number.

Nucleotide sequences were analyzed with the DNASIS program (Hitachi Software Engineering Co., Ltd., Yokohama, Japan). A homology search was carried out with databases of the GenBank, Swiss Plot, PDB and PIR by using the software basic local alignment search tool in the BLAST network service (the National Center for Biotechnology Information, Bethesda, MD). Phylogenetic analysis was performed by using the PHYLIP software package (version 3.5). An evolutional distance matrix, generated by using the Kimura formula in the PROTDIST, was used for construction of a phylogenetic tree by using the unweighted pair-group method analysis (UPGMA) (Felsenstein, J. 1989. PHYLIP-phylogeny inference package (version 3.3). Cladistics 5:164-166). The data were also examined using parsimony analysis (PROTPARS in PHYLIP). A bootstrap analysis was carried out to investigate the stability of randomly generated trees by using SEQBOOT and CONSENSE in the same package. The nucleotide sequence of the *p28* gene and its gene copies has been assigned GenBank accession numbers U72291 and AF021338, respectively.

Proteins of the E. chafeensis omp-1 Family.

Five complete omp-1 gene copies (omp-1B to omp-1F) encode 279 to 287-amino acid proteins with molecular masses of 30,320 - 31,508 Da. Omp-1A encodes an 82-amino acid partial protein (9,243 Da) which lacks the N-terminal region. The 25-amino acid sequence at the N-terminus of OMP-1B to OMP-1F (encoded in omp-1B to omp-1F) is predicted to be a signal peptide because three carboxyl-terminal amino acids of the signal peptides (Ser-X-Ala in OMP-1B, Leu-X-Ser for OMP-C, and Ser-X-Ser for OMP-1D and OMP-1F) are included in the preferred amino acid sequence of signal peptidase at the processing sites proposed by Oliver .. The calculated molecular masses of the mature OMP-1B to OMP-1F from the predicted amino acid sequences are 28,181 Da for OMP-1B, 27,581 Da for OMP-1C, 28,747 Da for OMP-1D, 27,776 Da for OMP-1E, and 27,933 Da for OMP-1F. The estimated isoelectric points are 4.76-5.76 in the mature OMP-1B to OMP-1F. An amino acid sequence in omp-1F gene (the 80th to 94th amino acids) was identical to the N-terminal amino acid sequences of E. chafeensis native P23 protein as determined chemically, which indicates that P23 is derived from the omp-1F gene. Amino acid sequences identical to the N-terminal sequences of P25, P27, and P29 were not found in those from omp-1 gene copies cloned in this study.

Alignment of predicted amino acid sequences of the *E. chafeensis* OMP-1 family and *Cowdria ruminantium*, revealed substitutions or deletions of one or several contiguous amino acid residues throughout the molecules. The significant differences in sequences among the aligned proteins are seen in the regions indicated SV (semivariable region) and HV (hypervariable region) 1 to 3 in Fig 31. Computer analysis for hydropathy revealed that protein molecules predicted from all *omp-1* gene copies contain alternative hydrophilic and hydrophobic motifs which are characteristic of transmembrane proteins. The HV1 and HV2 were found to locate in the hydrophilic regions.

The amino acid sequences of 5 mature proteins without signal peptides (OMP-1C to OMP-1F and a P28) were similar to one another (71-83%) but the sequence of OMP-1B was dissimilar to those of the 5 proteins (45-48%). The amino acid sequences of the 5 proteins showed an intermediate degree of similarity with that of C. ruminantium MAP-1 (59-63%), but the similarity between that of the OMP-1B and the C. ruminantium MAP-1 was low (45%). These relations are shown in a phylogenetic tree which was obtained based on the amino acid sequence alignment by UPGMA method in the PHYLIP software package (Fig. 10). Three proteins (P28, OMP-1D, and OMP-1F) and two proteins (OMP-1C and OMP-1E) formed two separate clusters. The OMP-1B was located distantly from these two clusters. The C. ruminantium MAP-1 was positioned between the OMP-1B and other members in the OMP-1 family.

Preparation of a Recombinant form of OMP-1 and P30

The 0.8-kb p28 gene was excised from the clone pCRIIp28 by EcoRI-NotI double-digestion, ligated into EcoRI-NotI sites of a pET 29a expression vector, and amplified in Escherichia coli BL21 (DE3)pLysS (Novagen, Inc., Madison, WI). The clone (designated pET29p28) produced a fusion protein with a 35-amino acid sequence

carried from the vector at the N terminus. The amino acid sequence of the OMP-1 portion of the fusion protein is depicted in Fig. 1.

An expression vector comprising the p30 gene was used to prepare the recombinant form of P30.

The following examples are for purposes of illustration only and are not intended to limit the scope of the claims which are appended hereto.

Preparation of anti rP28 (anti-OMP1) antibody

The (r) P28 antigen was prepared by excising the gel band corresponding to the rP28 in SDS-PAGE, mincing the band in phosphate-buffered saline (PBS), pH 7.4, and mixing with an equal volume of Freund's incomplete adjuvant (Sigma). The rP28 mixture (1 mg of protein each time) was subcutaneously injected into a rabbit every 2 weeks four times. A serum sample was collected from the rabbit to provide the anti-rP28 antibody

The anti-rP28 antibody was examined by western immunoblots analysis. The results indicated that the rabbit anti-rP28 antibody recognized not only rP28 (31 kDa) and P28, but also P29 and P25 of E. chafeensis and P30 of E. canis. These results indicate that P28 shares antigenic epitopes with P25 and P29 in E. chafeensis and P30 of E. canis.

Example 1. Assaying for the presence of anti-OMP-1 antibody in a Patient

Convalescent-phase serum from a patient with clinical signs of human ehrlichiosis was used. Western blot analyses using the rP28 protein as antigen was performed with 1:1,000 dilutions of this serum. Alkaline phosphatase-conjugated affinity-purified anti-human, anti-rabbit or anti-mouse immunoglobulin G (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD) were used at a 1:1,000 or 1:2,000 dilution as secondary antibodies. Results indicated that serum from a patient with clinical signs of human ehrlichiosis reacted strongly to rP28 (31 kDa).

Example 2 Assaying for the presence of anti-OMP-1 antibody in a Patient

Convalescent-phase serum from a patient with clinical signs of human ehrlichiosis was reacted with the rP30 protein of E.canis as described in Example 1. The serum reacted strongly to rP30. These results indicate the rP30 is useful for diagnosing an infection with E. chafeensis in human patients.

Example 3. Identifying E. chafeensis-infected cells using anti-rP 28 antibody

E. chafeensis-infected DH82 cells were sonicated and centrifuged at 400 x g for 10 min. The supernatant was then centrifuged at 10,000 x g for 10 min to obtain ehrlichia-enriched pellet. The pellet was resuspended and incubated with rabbit anti-rP28 antibody or normal rabbit serum (1:100 dilution) at 37°C for 1h in PBS containing 1% bovine serum albumin (BSA-PBS). After washing, the ehrlichiae was incubated with gold-conjugated protein G (20 nm), Sigma) at 1:30 dilution for 1 h at room temperature in BSA-PBS. After washing again, the specimen was fixed with 1.25% formaldehyde, 2.5% glutaraldehyde, and 0.03% trinitrophenol in 0.1 M cacodylate buffer (pH 7.4) for 24h and postfixed in 1% osmium-1.5% potassium ferricyanide for 1 h (34). The section was then embedded in

PolyBed 812 (Polysciences, Warraington, Pa). The specimen was ultrathin sectioned at 60 nm, stained with uranyl acetate and lead citrate, and observed with a Philips 300 transmission electron microscope at 60 kV.

Transmission immunoelectron microscopy with colloidal gold-conjugated protein G and rabbit anti-rP28 antibody revealed gold particles bound to *E. chafeensis* surface. The distribution of the particles was random, close to the surface, and appeared as if almost embedded in the membrane, suggesting that the antigenic epitope protrudes very little from the lipid bilayer. Nonetheless, the antigenic epitope was surface-exposed, and thus, could be recognized by rabbit anti-rP28 antibody. No gold particles were observed on host cytoplasmic membrane or *E. chafeensis* incubated with normal rabbit serum.

Example 4. Immunization of mice and E. chafeensis challenge.

The rP28 band in SDS- PAGE was excised, minced, and mixed with an equal volume of Freund's incomplete or complete adjuvant. Nine BALB/c male mice (6 weeks old) were divided into two groups. Five mice were intraperitoneally immunized a total of four times at 10-day intervals; twice with a mixture of the minced gel with the rP28 (30 to 40 µg of protein per mouse each time) and incomplete adjuvant, and twice with a mixture of the recombinant protein (the same amount as before) and complete adjuvant. Four mice were intraperitoneally injected with a mixture of the minced gel without protein and the respective adjuvants. For ehrlichia-challenge, approximately 1 x 10⁷ DH82 cells heavily-infected with E. chafeensis were disrupted by sonication in serum-free DMEM (GIBCO-BRL) and centrifuged at 200 x g for 5 min. The supernatant was diluted to a final volume of 5 ml, and 0.3 ml was inoculated intraperitoneally into each mouse 10 days after the last immunization. Before challenge, all 5-immunized mice had a titer of 1:160 against E. chafeensis antigen by IFA and all 4-nonimmunized mice were negative.

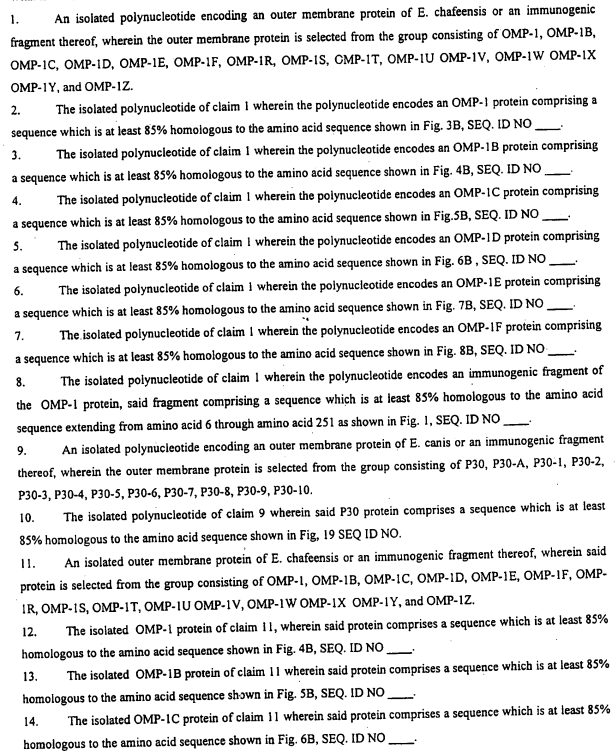
At day 5 post-challenge, approximately 1 ml of blood was collected in an EDTA tube from each mouse and protection was assessed by PCR detection of *E. chafeensis* 16S rDNA in the buffy coat of the collected blood. *E. chafeensis* could not be reisolated in cell culture at day 10 postinfection. Day 5 post challenge is the optimum time at which establishment of ehrlichial infection can be examined by PCR without the influence of residual DNA from the ehrlichiae used as the challenge before the spontaneous clearance of organisms take place. The *E. chafeensis*-specific DNA fragment was observed in all nonimmunized mice but not in any immunized mice, indicating that immunization of rP28 apparently protects mice from ehrlichial infection and indicating that the P28 is a potential protective antigen

Example 5 Assaying for the presence of anti-P30 antibody in Dogs

The rP30 protein was used as an antigen in a Western immunoblot analysis and dot blot analysis to detect the presence of antibody to E. canis in serum from E-canis infected dogs. The results of the Western immunoblot analysis indicated that reactivity of the sera with rP30 was stronger than the reactivity that was observed when purified E.canis was used as antigen. The results of the dot blot assay indicated that rP30 is a useful and sensitive tool for serodiagnosis of canine ehrlichiosis.

CLAIMS

What is claimed is:



•	
15.	The isolated OMP-1D protein of claim 11 wherein said protein comprises a sequence which is at least 85%
homolog	ous to the amino acid sequence shown in Fig. 7B, SEQ. ID NO
16.	The isolated OMP-1E protein of claim 11 wherein said protein comprises a sequence which is at least 85%
homolog	ous to the amino acid sequence shown in Fig. 8B, SEQ. ID NO
17.	The isolated OMP-1F protein of claim 11 wherein said protein comprises a sequence which is at least 85%
homolog	ous to the amino acid sequence shown in Fig. 9B, SEQ. ID NO
18.	The isolated immunogenic fragment of the OMP-1 protein of claim 11, said fragment comprising a
sequenc	e which is at least 85% homologous to the amino acid sequence extending from amino acid 6 through amino
acid 251	as shown in Fig. 1, SEQ. ID NO
19.	An isolated outer membrane protein of E. canis or an immunogenic fragment thereof, wherein the outer
membra	ne protein is selected from the group consisting of P30, P30-A, P30-1, P30-2, P30-3, P30-4, P30-5, P30-6,
P30-7, F	² 30-8, P30-9, P30-10.
20.	The isolated P-30 protein of claim 19 wherein said protein comprises a sequence which is at least 85%
homolo	gous to the amino acid sequence shown in Fig 19, SEQ ID NO
21.	A method for diagnosing an infection with E. chafeensis in a patient comprising the steps of:
	(a) providing a serum sample from the patient;
	(b) providing an outer membrane protein selected from the group consisting of a protein of claim
ll, a pr	otein of claim 19, and mixtures thereof;
	(c) contacting the serum sample with the outer membrane protein; and
	(d) assaying for the formation of a complex between antibodies in the serum sample and
	the outer membrane protein, wherein formation of said complex is
	indicative of infection with E. chafeensis.
22.	A method for diagnosing an infection with E. canis in a Canidae patient comprising the steps of:
	(a) providing a serum sample from the patient;
	(b) providing an outer membrane protein of claim 19;
	(c) contacting the serum sample with the outer membrane protein; and
	(d) assaying for the formation of a complex between antibodies in the serum sample and
	the outer membrane protein, wherein formation of said complex is
	indicative of infection with E. canis.
	\cdot

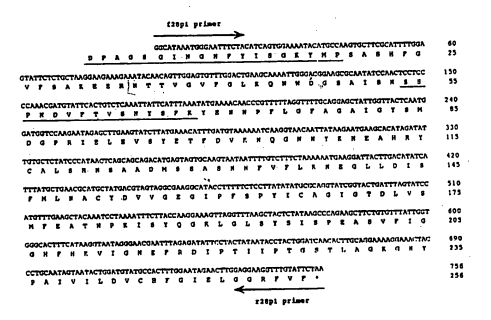


Fig. 1

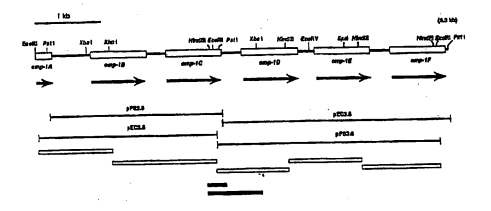


Fig. 2

60	50	40	30		10
TTCTCTACCT	CATTAATATC	GCATTGATAT	CATAACAAGT	AAAAAGTTTT	
. 120	110	100	90	80	70
CATCAGTGGA	GTAATTTCTA	GGTATTAACG	AGCAGGTAGT	TTTCCGACCC	GGAGTATCAT
180	170	160	150	140	130
AAGAAATACA	CTAAGGAAGA	GTATTCTCTG	GCATTTTGGA	CAAGTGCTTC	AAATACATGC
240	230	220	210	200	190
CAACTCCTCC	GCGCAATATC	TGGGACGGAA	GAAGCAAAAT	TGTTTGGACT	ACAGTTGGAG
300	290	280	270	260	250
CCCGTTTTTA	ATGAAAACAA	TCATTTAAAT	CTCAAATTAT	TATTCACTGT	CCAAACGATG
360	350	. 340	330	320	310
TGAAGTATCT	GAATAGAGCT	GATGGTCCAA	TTACTCAATG	GAGCTATTGG	GGTTTTGCAG
420	410	400	390	380	370
	AGAATGAAGC	AACAATTATA	AAATCAAGGT	TTGATGTAAA	TATGAAACAT
480	470	460	450	440	430
	CAAGTAATAA	ATGAGTAGTG	AGCAGCAGAC	CCCATAACTC	TGTGCTCTAT
540	530	520	510	500	490
TGACGTAGTA	ACGCATGCTA		TGACATATCA	AAGGATTACT	CTAAAAAATG
600	590	580	570	560	550
TTTAGTATCC		TGCGCAGGTA	TCCTTATATA	TACCTTTTTC	GGCGAAGGCA
660	650	640	630	620	610
	AGTTAGGTTT		TAAAATTTCT	CTACAAATCC	ATGTTTGAAG
720	710	700	690	680	670
AGGGAACGAA		GGGCACTTTC	GTTTATTGGT	AAGCTTCTGT	ATAAGCCCAG
780	770	760	750	740	730
	TTGCAGGAAA		AATACCTACT	TTCCTACTAT	TTTAGAGATA
840	830	820	810	800	790
	AACTTGGAGG			TAATACTGGA	CCTGCAATAG
900	890	880	870	860	850
300	. 690		••••	• • • • • • • • •	TTCTAA

Fig. 3A

60	50	40	30	20	. 10
VFSAKEERNT	KYMPSASHFG	GINGNFYISG	GVSFSDPAGS	ALISLISSLP	MNYKKVFITS
	110	100	90		· 70
DGPRIELEVS	GFAGAIGYSM	SFKYENNPFL	PNDVFTVSNY	WDGSAISNSS	TVGVFGLKQN
	· 170	160	150	140	130
FMLNACYDVV	LKNEGLLDIS	MSSASNNFVF	CALSHNSAAD	NNYKNEAHRY	YETFDVKNQG
	230	220	210	200	190
GHFHKVIGNE	ISPEASVFIG	YQGKLGLSYS	MFEATNPKIS	CAGIGTDLVS	GEGIPFSPYI
300	290	280	270	260	250
	F	FGIELGGREV	PAIVILDVCH	GSTLAGKGNY	FRDIPTILPT

Fig. 3B

					•
10	20	30	40	50	60
ATGAATTACA	AGAAAATTTT	TGTAAGCAGT	GCATTAATTT	CATTAATGTC	AATCTTACCT
70	80	90	100	110	. 120
TACCAATCTT	TTGCAGATCC	TGTAACTTCA	AATGATACAG	GAATCAACGA	CAGCAGAGAA
130	140	150	160	170	180
GGCTTCTACA	TTAGTGTAAA	GTATAATCCA	AGCATATCAC	ACTTCAGAAA	ATTCTCAGCT
190	200	210	220	230	240
GAAGAAGCTC	CCATCAATGG	AAATACTTCT		AGGTTTTCGG	GCTGAAAAA
250	260	270	280	290	300
GACGGAGATA	TAGCACAATC	TGCGAATTTT	AACAGGACAG	ATCCAGCCCT	CGAGTTTCAG
310	320	330	340	350	360
AATAACCTAA	TATCAGGATT	CTCAGGAAGT	ATTGGTTATG	CTATGGATGG	GCCAAGAATA
370	380	390	400	410	420
GAACTTGAAG	CTGCATACCA	AAAATTTGAT	GCAAAAAATC	CTGACAACAA	TGACACTAAT
430	440	450	460	470	480
AGCGGTGACT	ACTATAAATA	CTTTGGACTA	TCTCGTGAAG	ACGCAATAGC	AGATAAGAAA
490	500	510	520	530	540
TATGTTGTCC	TTAAAAATGA	AGGCATCACT	TTTATGTCAT	TAATGGTTAA	CACTTGCTAT
550	560	570	580	590	600
GACATTACAG	CTGAAGGAGT	ACCTTTCATA	CCGTATGCAT	GTGCAGGTGT	AGGAGCAGAC
610	620	630	640	650	660
CTTATAAACG	TATTTAAGGA	TTTTAATTTA	AAATTCTCAT	ACCAAGGGAA	AATAGGTATT
670	680	690	700	710	720
AGCTATCCAA	TCACACCAGA	AGTTTCCGCT	TTTATTGGAG	GATACTACCA	CGGAGTTATA
730	740	750			780
GGAAATAATT	TTAACAAAAT	ACCTGTAATA	ACACCTGTAG	TATTAGAAGG	AGCTCCTCAA
790	800	810			840
ACCACATCTG	CGCTAGTAAC	TATTGACACT	GGATACTTTG	GCGGAGAAGT	TGGAGTAAGG
850				890	900
TTCACCTTCT	AG	• • • • • • • • •	• • • • • • • • •		
		·	4.4		
	•	Fig	, 4A	•	
10	20	30	40	50	60
				GFYISVKYNP	SISHFRKFSA
70		90			120
FEX DINGNIE	TTKKVFGLKK	DGDTAOSANE			IGYAMDGPRI
	140			170	180
					FMSLMVNTCY
190				230	240
カエボカラ こうしょうしょう フェルファイン フェルファイン ファイン ファイン ファイン ファイン ファイン ファイン ファイン	PYACAGUGAD				FIGGYYHGVI
250				290	300
				FTF	
GMME WYTEAT	. TEAADEGNE	, TIGHTY TED			

Fig. 4B

		,			
10	2,0	30	40	50	60
ATGAACTGCA	AAAAATTTTTT	TATAACAACT	GCATTGGCAT	TGCCAATGTC	TTTCTTACCT
70	80	90	100	110	120
GGAATATTAC	TTTCTGAACC	AGTACAAGAT	GACAGTGTGA	GTGGCAATTT	CTATATTAGT
130	140	150	160	170	. 180
GGCAAGTACA	TGCCAAGTGC	TTCTCATTTT	GGAGTTTTCT	CTGCCAAAGA	AGAAAAAAAT
190	200	210	220	230	240
CCTACTGTCG	CGTTGTATGG	TTTGAAACAA	GATTGGAACG	GTGTTAGTGC	TTCAAGTCAT
250	260	270	280	290	300
GCTGATGCGG	ACTTTAATAA	CAAAGGTTAT	TCTTTTAAAT	ACGAAAACAA	
310	320	330	340	350	360
GGTTTTGCAG	GAGCTATTGG	TTATTCAATG	GGTGGTCCAA	GAATAGAGTT	TGAAGTGTCC
370	380	390	400	410	420
TATGAAACAT	TTGACGTGAA	AAATCAAGGT	GGTAATTACA	AAAATGATGC	TCACAGATAC
430	440	450	. 460	470	480
TGTGCCTTAG	ATCGTAAAGC	AAGCAGCACT	AATGCCACAG	CTAGTCACTA	CGTGCTACTA
490	500	510	520	530	540
ÄAAAATGAAG	GACTACTTGA	TATATCACTT	ATGTTGAATG	CATGCTATGA	CGTAGTAAGT
550	560	570	580	590	600
GAAGGAATAC	CTTTCTCTCC	TTACATATGT	GCAGGTGTTG	GTACCGATTT	AATATCCATG
610	620	630	640	650	660
TTTGAAGCTA	TAAACCCTAA	AATTTCTTAT	CAAGGAAAGT	TAGGTTTGAG	TTACTCTATA
670	680	690	700	710	720
AACCCAGAAG	CTTCTGTCTT	TGTTGGTGGA	CATTTTCATA	AAGTTGCAGG	
730	740	750	760	770	780
AGGGACATTT	CTACTCTTAA	AGCGTTTGCT	ACACCATCAT	CTGCAGCTAC	TCCAGACTTA
790	800	810	820	. 830	840
GCAACAGTAA	CACTGAGTGT	GTGTCACTTT	GGAGTAGAAC	TTGGAGGAAG	ATTTAACTTC
850	860	870	880	890	900
TAA					• • • • • • • • •
				•	
	•	Fig.	5A		•
. 10	20	30	40	50	60

. 10	20	30	40	50	60
MNCKKFFITT	ALALPMSFLP	GILLSEPVQD	DSVSGNFYIS	GKYMPSASHF	GVFSAKEEKN
70	80	90	100	110	120
PTVALYGLKQ	DWNGVSASSH	ADADFNNKGY	SFKYENNPFL	GFAGAIGYSM	GGPRIEFEVS
130	140	. 150	160	170	180
YETFDVKNQG	GNYKNDAHRY	CALDRKASST	NATASHYVLL	KNEGLLDISL	MLNACYDVVS
190	200	210	220	230	240
EGIPFSPYIC	AGVGTDLISM	FEAINPKISY	QGKLGLSYSI	NPEASVFVGG.	HEHKVAGNEF
250	260	270	280	290	300
RDISTLKAFA	TPSSAATPDL	ATVTLSVCHF	GVELGGRENE		

Fig. 5B

10		50	30	~ ~ ~	60
ATGAACTGCG			GCATTAACAT	TACTAATGTC	CTTCTTACCT
70	• •	50	100	110	120
GGAATATCAC	- 111010MICC	AGTACAGGAT	GACAACATTA	GTGGTAATTT	CTACATCAGT
130	210	150	160	170	180
GGAAAGTATA	TGCCAAGCGC	TTCGCATTTT	GGAGTTTTTT	CTGCCAAGGA	AGAAAGAAAT
190	200	210	220	230	240
ACAACAGTTG	Gagtatttgg	AATAGAGCAA	GATTGGGATA	GATGTGTAAT	ATCTAGAACC
250	260	270	280	290	300
ACTTTAAGCG	ATATATTCAC	CGTTCCAAAT	TATTCATTTA	AGTATGAAAA	
310	320	330	340	350	360
TCAGGATTTG	CAGGAGCTAT	TGGCTACTCA	ATGGATGGCC	CAAGAATAGA	GCTTGAAGTA
370	380	390	400	410	420
TCTTATGAAG	CATTCGATGT	TAAAAATCAA	GGTAACAATT	ATAAGAACGA	
430	440	450	460	470	480
TATTATGCTC	TGTCCCATCT	TCTCGGCACA	GAGACACAGA		AGGCAGTGCG
490	500	510	520	530	540
TCTGTCTTTC	TAATAAATGA	AGGACTACTT	GATAAATCAT	TTATGCTGAA	
550	560	570	580	590	600
GATGTAATAA	GTGAAGGCAT	ACCTTTTTCT	CCTTATATAT	GTGCAGGTAT	TGGTATTGAT
610	620	630	640	650	660
TTAGTATCCA	TGTTTGAAGC	TATAAATCCT	AAAATTTCTT	ATCAAGGAAA	ATTAGGCTTA
670	680	690	700	710	720
AGTTACCCTA	TAAGCCCAGA	AGCTTCTGTG	TTTATTGGTG	GACATTTTCA	TAAGGTGATA
730	740	750	760	770	780
GGAAACGAAT	TTAGAGATAT	TCCTACTATG	ATACCTAGTG		TGCAGGAAAA
790	800	810	820	830	840
GGAAACTACC	CTGCAATAGT	AACACTGGAC			
850	860	870	880	890	900
AGGTTTAACT	TCCAACTTTG	A	• • • • • • • • • • • • • • • • • • • •	090	900

Fig. 6A

10	20	30	40	50	60
MNCEKFFITT	ALTLLMSFLP	GISLSDPVQD	DNISGNFYIS	GKYMPSASHF	GVFSAKEERN
70	80	90	100	110	120
TTVGVFGIEQ	DWDRCVISRT	TLSDIFTVPN	YSFKYENNLF	SGFAGAIGYS	MDGPRIELEV
130	140	150	160	170	180
SYEAFDVKNQ	GNNYKNEAHR	YYALSHLLGT	ETQIDGAGSA	SVFLINEGLL	DKSFMLNACY
190	. 200	210	220	230	240
DVISEGIPFS	PYICAGIGID	LVSMFEAINP	KISYQGKLGL	SYPISPEASV	FIGGHFHKVI
250	260	270	280	290	300
GNEFRDIPTM	IPSESALAGK	GNYPAIVTLD	VFYFGIELGG	RFNFQL	

Fig. 6B

••	
10 20 30 40 50	60
ATGAATTGCA AAAAATTTTT TATAACAACT GCATTAGTAT CACTAATGTC CT	TTCTACCT
70 80 90 100 110	120
GGAATATCAT TTTCTGATCC AGTGCAAGGT GACAATATTA GTGGTAATTT CT	ATGTTAGT
130 140 150 160 170	180
GGCAAGTATA TGCCAAGTGC TTCGCATTTT GGCATGTTTT CTGCCAAAGA AG	TAAAAAAT
190 200 210 220 230	240
CCTACTGTTG CATTGTATGG CTTAAAACAA GATTGGGAAG GGATTAGCTC AT	CAAGTCAC
250 260 270 280 290	300
AATGATAATC ATTTCAATAA CAAGGGTTAT TCATTTAAAT ATGAAAATAA CC	
310 320 330 340 350	360
GGGTTTGCAG GAGCTATTGG TTATTCAATG GGTGGTCCAA GAGTAGAGTT TG	AAGTGTCC
370 380 390 400 410	420
TATGAAACAT TTGACGTTAA AAATCAGGGT AATAACTATA AAAATGATGC TC	
430 440 450 460 470	480
TGTGCTTTAG GTCAACAAGA CAACAGCGGA ATACCTAAAA CTAGTAAATA CG	מיייים איני מיייים אינים
490 500 510 520 530	540
AAAACCCAAC CAMMOOMMCA CAMAMOAMA	DAAATAAI
550 560 570 580 590	600
GAGAGCATAC CTTTGTCTCC TTACATATGT GCAGGTGTTG GTACTGATTT AA	
610 620 630 640 650	660
TTTCAACCON CAAROCCON ARCON ARC	ACTCTATA
670 680 690 700 710	720
AACCCAGAAG CTTCTGTATT TATTGGTGGA CATTTTCATA AGGTGATAGG AA	
730 740 750 760 770	780
ACCCACAMMO CMACMONALA ACCAMMONALA	TAGCAATA
790 800 810 820 830	840
GTAACACTAA GTGTATGTCA TTTTGGAATA GAACTTGGAG GAAGGTTTAA CT	

Fig. 7A

10	20	30	40	50	. 60
MNCKKFFITT	ALVSLMSFLP	GISFSDPVQG	DNISGNEYVS	GKYMPSASHF	GMFSAKEEKN
70	80	90	100	110	120
PTVALYGLKQ	DWEGISSSSH	NDNHFNNKGY	SFKYENNPFL	GFAGAIGYSM	GGPRVEFEVS
130	140	150	160	170	180
YETFDVKNQG	NNYKNDAHRY	CALGOODNSG	IPKTSKYVLL	KSEGLLDISF	MLNACYDIIN
190	200	210	220	230	240
ESIPLSPYIC	AGVGTDLISM	FEATNPKISY	QGKLGĻSYSI	NPEASVFIGG	HFHKVIGNEF
250	260	270	280	290	300
RDIPTLKAFV	TSSATPDLAI	VTLSVCHFGI	ELGGRENE.		

Fig. 7B

10		50	40	50	60
ATGAATTGCA		TATAACAACT	ACATTAGTAT	CGCTAATGTC	
70	•	J	100	110	120
GGAATATCAT	CIONIGC	AGTACAGAAC	GACAATGTTG	GTGGTAATTT	CTATATCAGT
130	4.0	150	160	170	180
GGGAAATATG	TACCAAGTGT	TTCACATTTT	GGCGTATTCT	CTGCTAAACA	
190	200	210	220	230	
ACAACAACCG	GAGTATTTGG	ATTAAAGCAA	GATTGGGATG	GCAGCACAAT	ATCTAAAAAT
250	260	270	280	290	300
TCTCCAGAAA	ATACATTTAA	CGTTCCAAAT	TATTCATTTA		TAATCCATTT
310	320	330	340	350	360
CTAGGTTTTG	CAGGAGCTGT	TGGTTATTTA	ATGAATGGTC		- • •
370	380	390	400	410	420
TCCTATGAAA	CATTTGATGT	GAAAAACCAG	GGTAATAACT	ATAAGAACGA	•
430	440	450	460	470	480
TATTATGCTT	TAACCCATAA	CAGTGGGGGA	AAGCTAAGCA	ATGCAGGTGA	TAAGTTTGTT
490	500	510	520	530	540
TTTCTAAAAA	ATGAAGGACT	ACTTGATATA	TCACTTATGT	TGAATGCATG	CTATGATGTA
550	560	570	580	590	600
ATAAGTGAAG	GAATACCTTT	CTCTCCTTAC	ATATGTGCAG	GTGTTGGTAC	TGATTTAATA
610	. 620	630	640	650	660
TCCATGTTTG	AAGCTATAAA	CCCTAAAATT	TCTTATCAAG	GAAAGTTAGG	TTTGAGTTAC
670	680	690	700	710	720
TCCATAAGCC	CAGAAGCTTC	TGTTTTTGTT		TTCATAAGGT	GATAGGGAAT
730	740	750	760	770	780
GAATTCAGAG	ATATTCCTGC	TATGATACCC	_	CTCTCACAGG	TAATCACTTT
790	800	810	820	830	840
ACTATAGTAA	CACTAAGTGT	ATGCCACTTT			GTTTAACTTT
850	860	870	880	890	
TAA		•••••	• • • • • • • • • • • • • • • • • • • •	090	900
				•••••	

Fig. 8A

60	50	40	30	20	. 10
GVFSAKQERN	GKYVPSVSHF	DNVGGNFYIS	GISFSDAVQN	TLVSLMSFLP	MNCKKFFITT
120	. 110	100	90	80	70
MNGPRIELEM	LGFAGAVGYL	YSFKYENNPF	SPENTFNVPN	DWDGSTISKN	TTTGVFGLKQ
180	170	160	150	140	130
SLMLNACYDV	FLKNEGLLDI	KLSNAGDKFV	YYALTHNSGG	GNNYKNDAHK	SYETFOVKNO
240	230	220	210	200	190
GGHFHKVIGN	SISPEASVFV	SYQGKLGLSY	SMFEAINPKI	ICAGVGTDLI	ISEGIPFSPY
300	290	280	270	260	250
		GVELGGRENE	TIVTLSVCHF	STSTLTGNHF	EFRDIPAMIP

Fig. 8B

10	20	30	40	50	60
ATGGAAAATC	TCATGAATAA	GAAAAACAAA	TTCTTTACAA	TAAGTACAGC	AATGGTATGC
. 70	80	90	100	110	120
TTATTGTTAT	TACCTGGTAT	ATCATTTTCA	GAAACTATAA	ACAACAGTGC	TAAAAAACAG
130	140	150	160	170	180
CCTGGGTTAT	ATATCAGTGG	GCAGTACAAA	CCTAGTGTTT	CAGTTTTTAG	TAATTTTTCA
190	200	210	220	230	240
GTAAAAGAAA	CTAATGTTCC	CACAAAGCAG	TTAATAGCAC	TTAAAAAAGA	CATTAATTCT
250	260	270	280	290	300
GTTGCAGTTG	GTAGTAATGC	TACTACAGGT	ATTAGCAATC	CAGGTAATTT	CACAATTCCT
310	320	330	340	350	360
TATACTGCAG	AATTTCAAGA	TAATGTTGCC	AATTTCAATG		TTACTCTTTT
370	380	390	400	410	420
CCTGATAGTC	TAAGAATTGA	AATAGAGGGA	TTTCATGAAA	AATTTGATGT	CAAAAACCCT
430	440	450	460	47.0	480
GGAGGTTACA	CACAAGTAAA	AGATGCGTAC	CGTTATTTTG	CACTAGCACG	TGATTTAAAA
490	500	510	520	530	540
GATGGCTTCT	TTGAACCTAA	AGCGGAAGAT	ACAGGTGTTT	ATCATACTGT	
550	560	570	580	590	600
GATGGATTAT	CTATTTTATC	TACTATGGTŤ	AACGTCTGTT	ACGATTTTTC	
610	620	630	640	650	660
TTACCAGTCT	TACCTTATAT	ATGTGCAGGT	ATGGGTATAA	ACGCCATAGA	
670	680	690	700	710	720
GCTTTACATG	TAAAATTTGC	TTACCAAGGC	AAACTAGGTA	TTAGCTATCA	
730	740	750	760	770	780
AAAGTAAATT	TATTCCTTGA	TGGGTATTAC	CATCAAGTAA	TAGGCAATCA	
790	800	810	820	•	840
TTAAACGTAA	ACCATGTTTA			AAGTCACATC	
850					-
ACACTTGACA	TTGCATACTT	TGGTGGCGAA	GTTGGAATAA	GATTCACATT	TTAA

Fig. 9A

Uø	50	40	∪د	۷	±0
TKQLIALKKD	NFSVKETNVP	QYKPSVSVFS	KKQPGLYISG	SFSETINNSA	MVCLLLLPGI
120	110	100	90	80	. 70
IEGFHEKFDV	YSFPDSLRIE	NVANFNGAVG	TIPYTAEFQD	TTGISNPGNF	INSVAVGSNA
180	170	160	150	140	130
TMVNVCYDFS	MKNDGLSILS	AEDTGVYHTV	DLKDGFFEPK	DAYRYFALAR	KNPGGYTQVK
240	230	220	210	200	190
GYYHQVIGNQ.	LFTKVNLFLD	YQGKLGISYQ	FFDALHVKFA	CAGMGINAIE	VDELPVLPYI
300	290	280	270	260	250
	• • • • • • • • • •	GGEVGIRFTF	AVATLDIAYF	TLKESPKVTS	FKNLNVNHVY

Fig. 9B

60	50	40		~~	10
TTTATCATTT	TCTTAGCATA	GGAGAATATA	TACTAGAGTG	AAGAAAAACT	ATGATATATA
	110	100	90	80	70
CAGCCTTGCT		GTAAATATTA	TCTAGTGCTG	CTTATATCTT	ATTCTTTCTA
180	170	160	150	140	130
AAAATTAATA	TTAGCACAAA	ATCTTTAACG	AAGAACTAAT	TCAGTCTACT	ATATGTGTTA
	230	220	210	200	190
CGGTAAACCG	GTTATTTGTA	AACATGAATT	TAAGTTTAGT	GTCGTGATAC	AAAGATAAAT
300	290	280	270	260	250
AAATTACACA	GAAACTTTCA	TCCTTTATTA	TGGAATATTT	AAATTTTTTA	TTAAATTTAC
360	350	340	330	320	310
サカカのこのこのであ			TAAATGCGGC	CTAATGATAG	CTAATAATTC
400	410	400	390	. 380	370
42U		GAGTACCGTA	TACTGGCAGT	CATATACACT	CTACATTATA
480	470	460	450	440	430
46U	ACCCTTCTCT		TAAATTACTT	TCTGTCAATG	GAAAACATTA
	530	520	510	500	490
540 CAGTAATGAA	CTAGAGAGTT		AATAATACCA	ATACTCTCGT	CATAATAAAA
600	500	580	570	560	550
7 UUG	ATGAGTGCTA	GAAAGTTCTT	AATAAATAAG	GGAATATATC	ATTCGAGTAA

Fig. 10A

60	50	40	30	20	10
IFNVSTKKLI	ICVISLLRTN	VNIIRYNSLA	ILSTYIFLVL	GEYILAYLSF	MIYKEKLTRV
120	110	100	90	80	70.
FYTTLWDNPA	LIIPNDSKCG	SFIRNFQNNT	LNLQIFYGIF	NMNCYLYGKP	KDKCRDTKFS
180	170	160	150	140	130
IPNAREFSNE	HNKNTLVIIP	INYNRSVLNQ	ENIICQCKLL	EYRNFFDILY	LHYTYTLTGS
240	230	220	210	200	190
				ESSYEC	IRVRNISINK

Fig. 10B

10	20	30	40	50	60
ATGAATAAAA	AAAACAAGTT	TATTATAGCT	ACAGCATTGG	TATATTTACT	GTCATTACCT
70	80	90	100	110	120
AGTGTATCGT	TTTCAGAGGT	TACAAACAGC	AGTATTAAAA		GTTATATATT
130	140	150	160	170	
AGTGGACAAT	ACAAACCAAG	TGTTTCTGTT			180
190	200			TCTCAATTAA	AGAAACTAAC
		210	220	230	240
ACTATCACAA	AAAATCTTAT	AGCGTTAAAA	AAAGATATTA	ACTCTCTTGA	AGTTAACGCC
250	260	270	280	290	300
GATGCTAGTC	AAGGTATTAG	TCATCCAGGA			AGCAGCATTT
310	320	330	340	350	360
GAAGATAATG	CTTTTAATTT	CAACGGTGCT	-		
370				TTACTGAAGG	TCTAAGGATT
	380	390	400	410	420
GAAATAGAAG	GTTCCTATGA	AGAATTTGAT	GCTGAAAACC	CTGGAGGTTA	TGGTCTAAAT
430	440	450	460	47.0	480
GATGCCTTTC	GGTACTTTGC	TTTAGCACGT	GATATGGAAA	GCAACAAGTT	CCTACCAAAA
490	500	510	520		
GCACAAAGCT				530	540
	CAC	• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •	• • • • • • • • • •	

Fig. 11A

10	20	30	40	50	60
MNKKNKFIIA	TALVYLLSLP	SVSFSEVTNS	SIKKHSGLYI	SGQYKPSVSV	FSSFSIKETN
70	80	90	100	110	120
TITKNLIALK	KDINSLEVNA	DASQGISHPG	NFTIPYIAAF	EDNAFNFNGA	IGYITEGLRI
130	140	150	160	170	180
EIEGSYEEFD	AENPGGYGLN	DAFRYFALAR	DMESNKFLPK	AOSS	

Fig. 11B

10	20	30	40	50	60
				ATAAATTATC	CATCGCATCT
70	80	90	100	110	120
ATTATGGTTA	ACACCTGCTA	TGATATTTCA	ATTAATAATA	CATCAATAGT	ACCGTATTTA
130	140	150	160	170	180
TGCACAGGCA	TTGGTGAAGA	TCTTGTAGGG	CTTTTTAATA	CAATACATTT	TAAACTTGCA
190	200	210	220	230	240
TATCAAGGGA	AAGTTGGAAT	GAGTTATTTG	ATAAATAACA	ATATCCTATT	ATTTTCTGAC
250	260	270	280	290	300
ATATATTATC	ATAAAGTCAT	GGGTAACAGA	TTTAAAAATT	TGTACATGCA	ATATGTAGCT
310	320	330	340	350	360
GATCCTAATA	TTTCTGAAGA	AACTATACCT	ATATTAGCAA	AACTTGATAT	TGGTTATTTT
370	380	390	400	410	420
GGAAGTGAAA	TTGGAATAAG	GTTTATGTTT	AACTAA		• • • • • • • • •

Fig. 12A

60	50	40	30	20	10
LENTIHEKLA	CTGIGEDLVG	INNTSIVPYL	IMVNTCYDIS	TTNNKLSIAS	SRIHDENYAI
120			90	80	70
ILAKLDIGYF	DPNISEETIP	FKNLYMQYVA	IYYHKVMGNR	INNNILLESD	YOGKVGMSYL
180	170	160	150	140	130
				N	GSETGTREME

Fig. 12B

		·			
10	20	30	40	50	60
ATGACAAAGA	AATTTAATTT	TGTAAATGTT	ATATTAACAT	TTTTGTTATT	TCTTTTCCCA
70	80	90	. 100	110	120
CTTAAGTCAT	TTACAACATA	TGCAAATAAT	AACACAATCA	CTCAAAAAGT	TGGATTGTAC
130	140	150	160	170	. 180
ATAAGTGGTC	AATATAAGCC	AAGTATTCCT	CATTTCAAGA	ATTTTTCAGT	AGAAGAAAAT
100	200	210	220	230	240
GACAAAGTAG	TAGATTTGAT	AGGTCTTACA	ACTGATGTTA	CATATATCAC	AGAACATATA 300
250	260	270	280	290	300
TTACGAGATA	ATACAAAATT	CAACACTCAT	TATATTGCAA	AGTTCAAGAA	CAATTTTATA 360
210	320	330	. 340	350	300
AATTTCAGCA	GTGCAATTGG	TTATTATTCT	GGGCAAGGAC	CAAGGTTAGA	420
270	380	390	400	410	420
TCTTATGGGG	ATTTTGATGT	TGTAAATTAT	AAAAATTATG	CAGTACAAGA	TGTTAATAGA 480
420	440	450	460	4/0	400
TATTTTGCTT	TAGTACGTGA	AAAAAATGGT	TCAAATTTCT	CTCCAAAACC	ACATGAAACT 540
490	500	510	520	530	4.00
AGTCAACCCT	CTGACAGTAA	TCCTAAAAAG	TCTTTTTATA	CTTTAATGAA	GAATAATGGG 600
EEO	560	570	580	590	. 600
GTATTTGTTG	CATCAGTAAT	AATCAACGGT	'`TGTTATGATI	TTTCTTTTAA	TAACACAACA 660
610	620	630	640		
ATATCACCTT	ACGTATGTAT	AGGAGTTGGA	A GGAGATTTI	TAGAGTTTTT	TGAAGTAATG
670	680	690	700) /10	720
CATATCAAGI	TTGCTTGCCA			ATCCAATATC	TCCCTCTATT 780
730	740	750	760		,
ACTATTTTT	CTGATGCACA	A TTATCACAA	G GTCATAAAT	ATAAATTTAA	CAACCTACAT
700	n : 800	n 810	B 824	ט פ	, 0.0
GTTAAGTAT			A CCTACCATT	A CCTCTGCAAC n 890	AGCCAAACTA 900
850	960	0 87		· -	,
AACATTGAA'	r attttggtg	g tgaagttgg	G ATGAGATTT	A TATTTTAA.	
		T7	124		
		Fig.	. 13A		
		_		n 51	n 60
10) 20	3(0 4	•	,
		LKSFTTYAN	N NTITOKVGL	n 11	P HFKNFSVEEN 120
70). 80	9	0 10		•
	r TDVTYITEH	I LRONTKENT	H YIAKEKNNE	n 17	S GQGPRLEIES
130	0 14	0 15	0 16		-
SYGDFDVVN	A KNAYAODAN	R YFALVREKN	G SNESPKPHE .0 22	U	K SFYTLMKNNG 0 240
1.9	0 20	U 21	U COPTEFFE	O. M. HIKFACOSK	V GISYPISPSI
VFVASVIIN	G CYDESENNT	T. ISPIVCIGV	70 GDEIEEEN	0 29	300
25	0 26	u morever o	'U . 20 10 : DITTELTE		G MRFIF
TIFADAHYH	K AINNKENNT	U AVIDIETV	in Ellinurur		

Fig. 13B

		•			
. 10	20	30	40	50	60
. 10 ATGAGÇAAAA	TATTTOKKK	TACAATAGGA	ACAGTACTTG	CATCTCTATT	ATCATTCTTA
•	0.0	90	100	170	• == -
. //	ころ4中中中で14日	TATAAATCAT	AATCATACAG	GAAATAACAC	TAGTGGTATA
	140	150	100	110	
130	CCCACTAGA	ACCAGGAGTA	TCCCATTTTA	GCAATTTCTC	AGTAAAAGAA
	~~~	חוכ	ZZU	270	
190	200 200 ATACA	ACTAGTAGGA	TATAAAAAAA	GTGCGTCTTC	TATCGATCCT
		270	280	230	
250	200 ፈን አ አ ርጥጥጥር ይ	AGGTCCATAT	ACTGTTACAT	TTCAAGATAA	TGCTGCTAGT
	200	. aan	340	200	
310		TTCTTACCCC	GAAAGTCTAA	GACTTGAACT	TGAAGGTTCT
	200	. วดก	400	410	
370		AGATCCTAAA	GACTACTCAG	CAAAAGATGC	TTTTAGGTTT
	440	. 450	400	, 41.0	
430	)	CTCTACTACT	GTTCCTGAT	CTCAAAAATA	TACAGTTATG
	E 0.1	. 516	1 521	, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
490		TAATCAATC	ATGATCAAT	GTTGTTATGA	CTATCTTTT
		~ 571	יום כי	, ,	
- 550	)	יים 'סדביהביה ה	r`'GCAGGTATT	G GTGAAGATTI	CATTGAATTT
		~ 67	n 54	u ••••	
610		O	T CAAGGAAAA	C TAGGTATTA	TTATTACTTC
TTTGATACT		^ 60	n 70	υ / 1.	
67	0 00	U TOCTOSTOS	G TACTATCAT	A GAGTTATAG	G GAATAAATTT 0 780
		75			•
73	0 74	., mcmmcmmac	A CTTGATGA	T TTCCTAAAG	C AACTTCTGCA 0 840
	~ /	no 81	n 82	20 83	0 840
79	0 80		er cereaage	G GAGTAAAGT	T TACATTTTAA 900
			70 81	30 89	900
85	10 8	60 8	. •		
	• • • • • • • • •	• • • • • • • • •		• • • • • • • • • • • • • • • • • • • •	
	•	r:	g. 14A		
rig. 14th					
	• .				
3	10	20			50 60
MSKKKFIT:	IG TVLASLLS	FL SIESFSAI	NH NHTGNNTS	GI YITGQYRPO	SV SHFSNFSVKE
	70	90	90 1	00 T	TO
TNVDTIOL	VG YKKSASSI	DP NTYSNFQG	PY TVTFQDNA	AS FSGAIGYS	YP ESLRLELEGS
•	an 1	40 1	50 1	.6U I	70
YEKFDVKD	PK DYSAKDAF	RF FALARNTS	TT VPDAQKYT	VM KNNGLSVA	SI MINGCYDLSF
	00 2	ስስ 2	210 2	20 2	30
NNLVVSPY	IC AGIGEDFI	EF FDTLHIKI	AY OGKLGISY	YF FPKINVFA	GG YYHRVIGNKF
. 2	50 2	260 2	270 2	280 4	90.

Fig. 14B

### 14/31

KNLNVNHVVT LDEFPKATSA VATLNVAYFG GEAGVKFTF.

				5.0	60
10	20	30	40	50	
ATGAGTGCTA				TAGTATGTTT	AGTGTCATAC 120
70	80	90	. 100	110	
TTACCTACTA	AATCTTTGTC				180
130	140	150	160	170	
CTATATGTCA	GTGGACAATA			TTAGTAATTT	TTCACTTAAA
190	200	210	220	230	240
GAAACTTATA	CTGACACTAA	AGAGTTATTA		AAGATATTAA	GTCTATTACA
250	260	270	280	290	300
GATATAACAA	CAAATAAAAA	ATTCAACATT	CCTTATAACA	CAAAATTTCA	AGATAATGCT
310	320	330	340	350	360
GTTAGCTTCA	GTGCAGCTGT	TGGATATATT	TCCCAAGACA	GTCCAAGGGT	TGAGGTAGAA
370	380	390	400	410	420
TGGTCTTATG	AAGAATTTGA	CGTTAAAAAT	CCTGGTAATT	ACGTAGTAAG	TGAAGCCTTC
430	440	450	460	47.0	480
AGGTATATTG	CTTTAGCAAG	AGGAATTGAT	AATCTTCAAA	AATATCCTGA	AACAAATAAG
490	500	510	520	530	540
татсттстта	TAAAGAACAA	TGGCTTATCT	GTCGCATCCA	TTATAATCAA	TGGCTGTTAT
550	560	570	580	590	600
СВТТТТТСТТ	TAAACAATTT	AAAAGTATCA	CCTTACATAT	GCGTAGGGTT	TGGTGGGGAC
610	620	630	640	650	660
דב בה ביד מידית דב בה ביד מידית	TTTTTAGTGC	TGTAAGTTTT	AAATTTGCTT	ATCAAGGTAA	GGTAGGTATC
670	680	690			720
א כייייאייריר איי	таттстстаа	TATGATTATA	TTTGCTGACG	GATATTACCA	TAAGGTCATA
730	740			770	780
יים מתיים היים ביים	ייייים ברביייי ייייים ברביייי			GTCTTAACAG	TCATCCTAAG
790				830	840
730	これででれてでする	י יירייים איינייין יירייים איינייין			TGGGTTAAAA
TCTACTTTIG 850			880	890	900
	• • • • • • • • • • • • • • • • • • • •		,		
TTTATATITT	AA	• • • • • • • • •	••••••		
		Fie	g. 15A		
		(	5. 10.1		
10	20	30	40	50	60
MCDKKKT:FTT	GSVLVCLVSY	LPTKSLSNLN	NINNNTKCTG	LYVSGQYKPT	VSHFSNFSLK.
70	90	90	Tuu	770	
ENVIDAKETT.	GLAKDIKSIT	DITTNKKFNI	PYNTKFQDNA	VSFSAAVGYI	SQDSPRVEVE 180
4 3 6	140	150	160	110	
MEALELUINI	PCNYVVSFAF	RYIALARGID	NLOKYPETNK	YVVIKNNGLS	VASIIINGCY 240
400	200	210	220	230	
חבות גוננו בספט בינת גוננו בספט	בטט בער ער ברנים	TIEFFSAVSF	KFAYQGKVGI	SYPLFSNMII	FADGYYHKVI
		270	280	290	200
250	VOUS NOUDE	STFAVATI.NV			
<b>GNKFNNTNA</b>	UA A 2 TU 2 ULL	OTPW4WITH!			

Fig. 15B

10	20	30	40	50	60
ATCACTAAAA	TATTTTAAAA	TACAATAGGA	GCAACACTTA	TTCATATGTT	GTTACCTAAC
70	80	90	100	110	. 120
איים יים יים יים יים יים יים יים יים יים	CAGAAACTAT	TAACAATAAC	ACTGATAAAC	TTTCTGGGTT	ATATATAAGT
120	140	150	160	170	100
CCCLATATA	AACCAGGGAT	TTCTCATTTC	AGCAAATTTT	CAGTCAAAGA	AATCTATAAT
100	200	210	220	230	. 240
CATAACATTC	AACTAATTGG	GTTAAGACAC	AACGCAATTT	CTACTAGTAC	CCTTAATATT
250	260	270	280	290	300
AATACAGATT	TTAATATCCC	CTATAAAGTA	ACATTTCAAA	ATAACATTAC	CAGCTTTAGT
210	320	330	340	. 350	300
CCACCTATTG	GTTATTCTGA	TCCCACAGGG	GCAAGATTTG	AGCTTGAAGG	TTCTTATGAA
270	380	. 390	400	410	420
GAATTTGATG	TGACAGATCC	TGGAGACTGC	TTAATAAAAG	ATACCTATAG	ATATTTCGCT
430	440	450	460	4/0	400
TTAGCTAGAA	ACCCATCAGG	TTCTAGCCCT	ACCTCAAACA	ACTATACTGT	TATGAGAAAT
490	500	510	) 520	) 530	340
СУТССТСТТТ	CCATTACTIC	TGTTATATTI	AATGGCTGTT	ATGACATCTT	TTTAAAGGAT
E E A	560	570	580	) 590	800
TTAGAAGTAT	CACCTTATG	ATGTGTTGGT	r Gtaggtgga	ATTTTATAGA	ATTTTTTGAC
617	620	n 630	640	) 65(	, 660
GCATTACAC	TTAAATTAG	CATACCAAGG	C AAGTTAGGT	A TCAATTATCA	CTTATCGACT
<i>(</i> 7)	. 69	n 690	0 70	0 /1(	, ,20
CARGCARGO	G TATTTATTG	A TGGATATTA	T CATAAGGTT	A TAGGAAATCI	A ATTCAACAAT
72	n 74	ი 75	0 76	0 77	, ,
CTABATGTT	C AACACGTGG	C TAGTACAGA	T TTTGGACCT	G TATACGCAG'	AGCCACACTT 840
70	n 80	0 81	.0 82	0 83	0 040
AACATTGGT	T ATTTTGGTG	G TGAAATCGG	A ATTAGACTT	A CATTITAA.	
					•
			Fig. 16A		
				_	ი 60
1	.0 2	20 3	30 4		<b>V</b>
MSKKNFITI	G ATLIHMLLE	N ISFPETIN	N TDKLSGLYI	S GOYKPGISH	F SKFSVKEIYN 0 120
_	-	90	ลก บ	JU . ——	
DNIOLIGLE	H NAISTSTL	NI NTDFNIPY	KV TFQNNITS!	S GAIGYSDPT	G ARFELEGSYE
	1	40 . 1	50 LI	י בי טב	•
EFDVTDPGI	DC LIKDTYRY	FA LARNPSGS	SP TSNNYTVM	RN DGVSITSVI	F NGCYDIFLKD
<b>.</b>		^^ 7	1 N 2	2U <b>-</b> -	, •
LEVSPYVC	VG VGGDFIEF	FD ALHIKLAY	QG KLGINYHL	ST QASVFIDG	Y HKVIGNQFNN 90 300
	50 . 2	60 2	70 2	8U <del>*</del> -	-
* 17701777 C	mn ECDUVAUA	TT. NIGYEGGE	IG IRLTF		•• . • • • • • • • • •

Fig. 16B

LNVQHVASTD FGPVYAVATL NIGYFGGEIG IRLTF.....

) 60	50	40	30	) <u>2</u> 0	. 10
OU CTTATTATTC	でなってなってなる。	GGTGCATCAT	TTTTATAATA	GAAAAAGTTI	ATGAATAATA
_	110	100	90	80	70
	CTTATTTTAA	AGTAACCATA	AGGAAATGTA	CCTCTTCTAC	ACATCTGAGG
	170			140	130
	GCAAATTTTC		ACCAGGAGTT	GACAATATAG	TATATCAGTG
	000	220	210		190
CATAGGGAAC	ACATCAGTGT	CTTAAAAAGG	ACTAGTTGGG	ATACTACTCA	ACCAACTACA
300	290	280	270	260	250
TCAAGACAAT		TTTCCTTACA	AAATTTCAAC	CAACCTACAC	
360	350	340	330	320	310
			AATTGGATAC	TCAGTGGGGC	GCCATAAGTT
AATTGAAGTA	410	400	390	380	370
72722222	GATCTCCTAC	AATCCAGAAG	TGATGTTAAA	ATGAAGAATT	GAGGCTTCTT
AGACGCATAC 480	47.0	460	450	440	430
TGATGACACA		GGCACTAATA	TGCTATGGAT	CACTAGCACG	AGGTATTTTG
540	530	520	510	500	490
GATAAATGGG		TTATCAATTT	AAATGACGGG	CTGTCATGAG	AGAAAATTCA
600	590	580	570	560	550
	ATGTATGCGC	GTAGTACCGT	TGATATACCA'	TTACATTAGA	TGTTACAATT
660	650	640	630	620	610
	TTGCTCATCA		TAATGATTTA	TAGAGTTTTT	GGAGATTTCA
720	710	700	690	680	670
TTACCATAAA			CCCTGAAGTA	ATTCTATATC	GGTATTAGTT
780	770	760	750	740	730
AAGTGACGCT	・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・・		AAACTTACAC	ACAGATTTAA	GTAACAGGTA
AAGIGACGCT 840	830	820	810	800	790
CCARAMMCCA			TGCTACACTC	CATCTGCAGT	CCTAAGTTCA
900	890	880	870	860	850
900	690	•••••	• • • • • • • • • •	TATTTTAA	GTAAGATTTA

# Fig. 17A

• •					
10	20	30	40	50	60
MNNRKSFFII	GASLLASLLF	TSEASSTON			00
70		100001014	SNHTYFKPRL	YISGQYRPGV	SHFSKFSVKE
, , , ,	80	90	100	110	
TNYNTTQLVG	LKKDISVIGN	SNITTYTNEN	FDYTAFFORM	A TOBOON TON	
130	140	4.000.000.000		MISISGAIGY	LYSENFRIEV
		150	160	170	180
EASYEEFDVK	NPEGSATDAY	RYFALARAMD	GTNKSSPDDT	DEFTIMENT	TOTOGUNETUR
190	200	210		WE I AUMUDG	PRIRZAMING
			220	230	240
CYNFTLDDIP	VVPYVCAGIG	GDFIEFFNDL	HVKFAHOGKV	GTSYSTSDEV	ST ET NOVVUE
250	260	270			
THE CATE COMMENT OF		270	280	<b>290</b> .	300
VTGNRFKNLH	VUHVSDLSDA	PKFTSAVATL.	NVGYFGGEIG	VRFIF	

Fig. 17B

				,	
10	20	30	40	50	60
TAGCAGCACT	AAAAAACAGT	TTGGGTTATA	TGTTAGTGGA	CAACACCAGC	CTAGTGTTTC
70	80	90	100	110	120
TATTTTTAGC	AATTTCTCAG	TAAAGGAAAC	TAATTTTCCT	ACAAAGTATT	CTAGCAGCTT
130	140	150	160	170	180
CTTAAAAAAA	GACATTAATT	CTGTCGAATT	TGACGATAGT	GTTACTGCTG	GCATTAGTTA
190	200	210	220	230	240
CCCACTTAAT	TTCAGTACTC	CTTATATAGC	TGTATTTCAA	GATAATATTT	CTAATTTTAA
250	260	270	280	290	300
TGGCGCTATT	GGGTACACTT	TTGTTGAAGG	CCCAAGAATT	GAAATAGAAG	GTTCTTATGA
310	320	330	. 340	350	360
AGAATTCGAT	GTCAAAGACC	CTGGAAGATA	TACAGAAATA	CAAGATGCAT	ACCGTTACTT
370	380	390	400	410	420
TGCTTTAGCA	CGTGATATAG	ACTCTATTCC	TACTAGCCCA	AAAAATAGAA	CTTCACATGA
430	440	450	460	470	480
TGGCAACAGT	TCATATAAGG	TATACCACAC	TGTAATGAAA	AATGAAGGAC	TATCTATAAT
490	500	510	520	530	.540
ATCCATTATG	GTCAATGGCT	GCTATGATTT	TTCTTCAGAT	AATTTATCAA	TATTACCTTA
550	560	570	580	590	600
TGTATGTGGT	GGTATAGGTG	TAAATGCTAÌ	AGAGTTTTTC	GATGCATTAC	ATGTTAAATT
610	620	630	640	650	660
CGCGTGTCAG	GGTAAATTAG	GTATTACTTA	TCCATTATCT	TCCAACGTTA	GTTTATTTGC
670	680	690	700	710	720
TGGTGGATAT	TATCACCAAG	TAATGGGCAA	CCAATTTAAA	AATCTAAATG	TTCAACATGT
730	740	750	760	770	780
AGCTGAACTT	AATGACGCAC	CCAAAGTTAC	ATCTGCAGTA	GCTACACTTG	ACATTGGGTA
790	800	810	820	830	840
TTTTGGTGGT	GAAATTGGAG	CAAGGCTTAT	ATTTTAA		
,					

# Fig. 18A

10	20	30	40	50	60
SSTKKUEGI.Y	VSGQHQPSVS	IFSNFSVKET	NEPTKYSSSE	LKKDINSVEF	DDSVTAGISY
70	80	90	. 100	110	120
PLNFSTPYIA	VFQDNISNFN	GAIGYTFVEG	PRIEIEGSYE	EFDVKDPGRY	TEIQDAYRYF
130	140	150	160	170	180
ALARDIDSIP	TSPKNRTSHD	GNSSYKVYHT	VMKNEGLSII	SIMVNGCYDF	SSDNLSILPY
190	200	210	220	230	240
VCGGIGVNAI	EFFDALHVKF	ACQGKLGITY	PLSSNVSLFA	GGYYHQVMGN	<b>QEKNLNVQHV</b>
250	260	270	280	290	300
AELNDAPKVT	SAVATLDIGY	FGGEIGARLI	F		· • • • • • • • • • • • • • • • • • • •

Fig. 18B

		·			
10	20	30	40	50	60
ATGAATTGCA	AAAGATTTTT	CATAGCAAGT	GCATTGATAT.	CACTAATGTC	TTTCTTACCT
70	80	90	100	110	120
AGCGTATCTT	TTTCTGAATC	AATACATGAA	GATAATATAA	ATGGTAACTT	TTACATTAGT
130	140	150	160	170	180
GCAAAGTATA	TGCCAAGTGC	CTCACACTTT	GGCGTATTTT	CAGTTAAAGA	AGAGAAAAAC
190	200	210	220	230	240
ACAACAACTG	GAGTTTTCGG	ATTAAAACAA	GATTGGGACG	GAGCAACAAT	AAAGGATGCA
250	260	270	280	290	300
AGCAGCAGCC	ACACAATAGA	CCCAAGTACA	ATATTCTCCA	TTTCAAATTA	
310	320	330	340	350	360
TATGAAAACA	ATCCATTTTT	AGGGTTTGCA	GGAGCTATTG	GCTACTCAAT	GGGTGGTCCA
370	380	390	400	410	420
AGGGTAGAGT	TTGAAGTGTC	TTACGAAATA	TTTGATGTAA	AAAACCAAGG	
430	440	450	460	470	480
AAGAACGATG	CTCACAAATA	TTGCGCTTTA	TCAAGACACA	CCGGAGGTAT	
490	500	510	520	530	540
GGTCATCAAA	ATAAATTTGT	CTTCCTAAAA	AATGAAGGAT		
550	560	570	580	590	600
ATAAACGCAT	GTTATGATAT	AACAATCGAC'	AGCATGCCAT		
610	620	630	640	650	660
GGTATTGGTA	GTGACTTAGT	TTCGATGTTT			
670	680	690	700	710	720
GGAAAATTAG	GTGTAAGTTA	CTCCATAAGC	CCAGAAGCAT		
730		750	760		780
TTTCACAGAG	TTATAGGTAA	TGAATTTAAA			
790					840
ACAGAAATTA	AAGGCACACA	GTTTACAACA			
850		,			•
GAGCTTGGAG	GCAGGTTTAC	TTTTTAA	• • • • • • • • •	• • • • • • • • •	• • • • • • • • •
		,	101		
		Fı	g. 19A		
. 10	20	30	40	50	60
MNCKRFFIAS	ALISLMSFLP	SVSFSESIHE	DNINGNFYIS	AKYMPSASHF	GVFSVKEEKN
` 70		90			

60	50	40	30	20	. 10
GVFSVKEEKN	AKYMPSASHF	DNINGNFYIS	SVSFSESIHE	ALISLMSFLP	MNCKRFFIAS
120	110	100	90	80	70
GAIGYSMGGP	YENNPFLGFA	IFSISNYSFK	SSSHTIDPST	DWDGATIKDA	TTTGVFGLKQ
180	170	. 160	150	140	130
NEGLLDISLM	GHONKEVELK	SRHTGGMPQA	KNDAHKYCAL	FDVKNQGNSY	RVEFEVSYEI
240	230	220	210	200	190
PEASVEVGGH	GKLGVSYSIS	ETTNPKISYQ	GIGSDLVSMF	SMPFSPYICA	INACYDITID
300	290.	280	270	260	250
	ELGGRFTF	VTLNICHFGL	TEIKGTQFTT	DIPAITPAGA	FHRVIGNEFK

Fig. 19B

10	20	· 30	40	50	60
ATGAAATATA	AAAAAACTTT	TACAGTAACT	GCATTAGTAT	TATTAACTTC	CTTTACACAT
70	80	90	100.	110	120
TTTATACCTT	TTTATAGTCC	AGCACGTGCC	AGTACAATTC	ACAACTTCTA	
130	140	150	160	170	180
AAATATATGC	CAACAGCGTC	ACATTTTGGA	ATTTTTTCAG	CTAAAGAAGA	ACAAAGTTTT
190	200	210	220	230	240
ACTAAGGTAT	TAGTTGGGTT	AGATCAACGA	TTATCACATA	ATATTATAAA	CAATAATGAT
250	260	270	280	290	300
ACAGCAAAGA	GTCTTAAGGT	TCAAAATTAT	TCATTTAAAT	ACAAAAATAA	CCCATTTCTA
310	320	330	. 340	- 350	.360
GGATTTGCAA	GAGCTATTGG	TTATTCAATA	GGCAATTCAA	GAATAGAACT	AGAAGTATCA
370	380	390	400	410	420
CATGAAATAT	TTGATACTAA	AAACCCAGGA	AACAATTATT	TAAATGACTC	
430	440	450	460	470	`480
TGCGCTTTAT	CTCATGGAAG	TCACATATGC	AGTGATGGAA	ATAGCGGAGA	
490	500	510	520	530	540
GCAAAAACTG	ATAAGTTTGT	ACTTCTGAAA	AATGAAGGTT	TACTTGACGT	CTCATTTATG
550	560	570	580	590	600
TTAAACGCAT	GTTATGACAT	AACAACTGAA	AAAATGCCTT	TTTCACCTTA	
610	620	630	640	650	660
GGTATTGGTA	CTGATCTCAT	ATCTATGTTT	GAGACAACAC	AAAACAAAAT	
670	680	690	. 700		720
GGAAAGTTAG	GTTTAAACTA	TACTATAAAC		CTGTTTTTGC	AGGTGGGCAC
730	740	750	760		780
TTTCATAAAG	TAATAGGTAA			CTCTATTACC	
790	800				
AACATTAAAG				TGTGCCATTT	
850			880	890	300
ATTGGAAGTA	GATTTTTCTT	TTAA	• • • • • • • • •	• • • • • • • • •	• • • • • • • • • •

#### Fig. 20A

60
)SF
L20
EVS
180
SEM
240
GGH 300
300
• • •

Fig. 20B

					60
10	20 .	30	40 -		• •
አጥርጥጥጥልጥል	CTAATATATA	TATTCTGGCT	TGTATTTACT	TTGCACTTCC	ACTATTGTTA
70	. 80	90	100	110	. 120
שייייייייייייייייייייייייייייייייייייי	ACTATTTTAG	GTGTAATATG	AATTGCAAAA	AAATTCTTAT	AACAACTGCA
130	140	150	160	. 170	. 160
<b>サムつ</b> かなかなかか	TAATGTACTC	TATTCCAAGC	ATATCTTTTT	CTGATACTAT	ACAAGATGGT
190	200	210	220	230	240
AACATEGETG	GTAACTTCTA	TATTAGTGGA	AAGTATGTAC	CAAGTGTCTC	ACATTTTGGT
250	260	270 °	280	290	300
ACCTTCTCAG	CTAAAGAAGA	AAGCAAATCA	ACTGTTGGAG	TTTTTGGATT	AAAACATGAT
310	320	330	340	350	. 300
TECENTEGAN	GTCCAATACT	TAAGAATAAA	CACGCTGACT	TTACTGTTCC	AAACTATTCG
370	380	390	400	410	420
TTCTCTTTCG	AGAACAATCC	ATTTCTAGGG	TTTGCAGGAG	CTATCGGTTA	CTCAATGGGT
. 430	440	450	460	470	400
CCCCTAGAA	TAGAATTCGA	AATATCTTAT	GAAGCATTCG	ACGTAAAAAG	TCCTAATATC
490	500	510	520	530	,540
0CF 4407747774	ATGACGCGCA	CAGGTACTGC	GCTCTATCTC	ATCACACATC	GGCAGCCATG
550	560	570	580	590	600
CARCCTGATA	AATTTGTCTT	CTTAAAAAÄC	GAAGGGTTAA	TTGACATATC	ACTTGCAATA
610	620	630	640	650	uga
AATGCATGTT	ATGATATAAT	AAATGACAAA	GTACCTGTTT	CTCCTTATAT	ATGCGCAGGT
670	680	690	700	710	120
7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	ATTTGATTTC	TATGTTTGAA	GCTACAAGTC	CTARARTTTC	CTACCAAGGA
720	740	750	760	770	, ,,,,
AAACTGGGCI	TTAGTTACTC	TATTAATCC	GAAACCTCTG	TTTTCATCGG	TGGGCATTTC
. 700	800	810	820	830	040
CACAGGATCI	TAGGTAATGA	GTTTAGAGAI	ATTCCTGCA	TAGTACCTAG	TAACTCAACT
95	3 860	870	) 880	) 890	, 500
ACAATAAGT(	GACCACAATI	TGCAACAGT	A ACACTAAATO	TGTGTCACT	TGGTTTAGAA
910	920	930	940	) 951	, 500
CTTGGAGGA	A GATTTAACTT	CTAA			
	F	rig. 21A			
					60
10	20	30	3 40		60
MEVTNIYIL	A CIYFALPLLI	IYFHYFRCM	M NCKKILITT	LISLMYSIPS	ISFSDTIQDG
	^/	• 0/	n 1131	1	,
NMGGNEYIS	G KYVPSVSHE	S SFSAKEESK	S TVGVFGLKH	D WDGSPILKNI	HADFTVPNYS
	4 4 4 4	1 1 16	n 161		
FRYENNPFL	G FAGAIGYSM	GPRIEFEIS	Y EAFDVKSPN	I NYQNDAHRY	C ALSHHTSAAM 0 240
	200	דכ ח	n 22	U	•
EADKEVELK	M FCT.TDTSTA	I NACYDIIND	K VPVSPYICA	G IGTDLISME	E ATSPKISYQG 0 300
	- 26	n . 77	n 28	U 44	<b>U</b> .
KLGISYSIN	P ETSVFIGGH	F HRIIGNEFR	D IPAIVPSNS	T TISGPOFAT	A LITMACHE CITE
31		0 33	34	0 35	U. 300
					• • • • • • • • • •

Fig. 21B

					cò
10	20		40	50	60
ATGAATTGCA	AAAAAATTCT				CTATGCTCCA
70	80	90	100	110	120
AGCATATCTT	TTTCTGATAC				CATCAGTGGA
130	140	150	160	170	180
AAATATGTAC		ACATTTTGGT	GTTTTCTCAG	CTAAAGAAGA	AAGAAACTCA
190	200	210	220	230	240
ACTGTTGGAG	TTTTTGGATT	AAAACATGAT	TGGAATGGAG	GTACAATATC	TAACTCTTCT
250	260	270	280	290	300
CCAGAAAATA	TATTCACAGT	TCAAAATTAT	TCGTTTAAAT	ACGAAAACAA	CCCATTCTTA
310	320	330	. 340	350	360
GGGTTTGCAG	GAGCTATTGG	TTATTCAATG	GGTGGCCCAA	GAATAGAACT	TGAAGTTCTG
370	380	390	400	410	420
TACGAGACAT	TCGATGTGAA	AAATCAGAAC	AATAATTATA	AGAACGGCGC	ACACAGATAC
430	440	450	460	470	480
TGTGCTTTAT	CTCATCATAG	TTCAGCAACA	AACATGTCCT	CCGCAAGTAA	CAAATTTGTT
490	500	510	520	530	540
TTCTTAAAAA	ATGAAGGGTT	AATTGACTTA	TCATTTATGA	TAAATGCATG	CTATGACATA
550	560	570	580	590	600
ATAATTGAAG	GAATGCCTTT	TTCACCTTAT	ATTTGTGCAG	GTGTTGGTAC	TGATGTTGTT
610	620	630	640	650	660
TCCATGTTTG	AAGCTATAAA	TCCTAAAATT	TCTTACCAAG	GAAAACTAGG	ATTAGGTTAT
670	680	690	700	710	720
AGTATAAGTT	CAGAAGCCTC	TGTTTTTATC	GGTGGACACT	TTCACAGAGT	CATAGGTAAT
730	740	750	760	770	780
GAATTTAGAG	ACATCCCTGC	TATGGTTCCT	AGTGGATCAA	ATCTTCCAGA	AAACCAATTT
790	800	810	820	830	840
GCAATAGTAA	CACTAAATGT	GTGTCACTTT	GGTTTAGAAC	TTGGAGGAAG	ATTTAACTTC
850					900
TGA				• • • • • • • • •	• • • • • • • • •
		Fig	. 22A		
10	20	30	40	50	. 60
MNCKKILITT	ALMSLMYYAP	SISFSDTIQD	DNTGSFYISG	KYVPSVSHFG	VFSAKEERNS
70	80				
TVGVFGLKHD	WNGGTISNSS	PENIFTVQNY	SFKYENNPFL	GFAGAIGYSM	GGPRIELEVL
130	140	150	160	170	180
YETFDVKNQN	NNYKNGAHRY	CALSHHSSAT	nmssasnkfv		SFMINACYDI
190	200	210	220	230	240
IIEGMPFSPY	ICAGVCTDVV	SMFEAINPKI	SYQGKLGLGY	SISSEASVFI	GGHFHRVIGN
250	•				
EFRDIPAMVP	SGSNLPENQF	AIVTLNVCHF	GLELGGRFNF		• • • • • • • • •
			•		

Fig. 22B

PCT/US98/19600 WO 99/13720

10	20	30	40	50	60			
ATGAATTGTA	AAAAAGTTTT	CACAATAAGT	GCATTGATAT	CATCCATATA	CTTCCTACCT			
. 70	80	90	100	110	120			
AATGTCTCAT	ACTCTAACCC	AGTATATGGT	AACAGTATGT	ATGGTAATTT	TTACATATCA			
130	140	150	160	170	180			
GGAAAGTACA	TGCCAAGTGT	TCCTCATTTT		CAGCTGAAGA	AGAGAAAAA			
190	200	210	220	230	240			
AAGACAACTG	TAGTATATGG	CTTAAAAGGA		GAGATGCAAT	ATCTAGTCAA			
250	260	270	280	290	300			
AGTCCAGATG	ATAATTTTAC		TACTCATTCA	AGTATGCAAG	CAACAAGTTT			
310	320	330	340	350	360			
TTAGGGTTTG	CAGTAGCTAT	TGGTTACTCG		CAAGAATAGA	AGTTGAGATG			
370	380	390	400	410	420			
TCTTATGAAG	CATTTGATGT	GAAAAATCCA	GGTGATAATT	ACAAAAACGG	TGCTTACAGG			
430	440	450	460	470	480			
TATTGTGCTT	TATCTCATCA	AGATGATGCG	GATGATGACA	TGACTAGTGC	AACTGACAAA			
490	500	510	520	530	540			
TTTGTATATT	TAATTAATGA	AGGATTACTT		TTATGACAAA	CATATGTTAT			
550	560	570			600			
GAAACAGCAA	GCAAAAATAT	ACCTCTCTCT		GTGCAGGTAT	TGGTACTGAT			
610	620	630						
TTAATTCACA	TGTTTGAAAC	TACACATCCT	AAAATTTCTI	ATCAAGGAAA	GCTAGGGTTG			
670	680	690	700	710	120			
GCCTACTTCG	TAAGTGCAGA	GTCTTCGGTT	TCTTTTGGT	TATATTTTCA	TAAAATTATA			
730	740			<u> </u>				
AATAATAAGI	TTAAAAATGT	TCCAGCCATO			GATAGTAGGA			
790	800	810			•			
CCACAGTTTC	CAACAGTAAC	ATTAAATGT	A TGCTACTTT	GATTAGAACT	TGGATGTAGG			
850				990	900			
TTCAACTTC:	r AA		• • • • • • • • • • • • • • • • • • • •	• • • • • • • • •				
	Fig. 23A							

60	50	40	30	20	ΙU
GIFSAEEEKK	GKYMPSVPHF	NSMYGNFYIS			MNCKKVFTIS
120	110	100	90	80	70
IGSPRIEVEM	LGFAVAIGYS	YSFKYASNKF	SPDDNFTIRN	KLAGDAISSO	KTTVVYGLKG
190	170	. 160	150	140	130
NISFMINICY	FVYLINEGLL	DDDMTSATDK	YCALSHODDA	GDNYKNGAYR	SYEAFDVKNP
240	230	220	. ~ 210	200	190
SFGIYFHKII	AYFVSAESSV	KISYQGKLGL	LIHMFETTHP	PYICAGIGTD	ETASKNIPLS
300	290.		270	260	250
	FNF	CYFGLELGCR	POFATVILNY	VPTNSDETVG	

Fig. 23B

. 10	20	30	40	50	60
	AAAATTTCT	TATAACAACT	ACATTGGTAT	CACTAACAAT	TCTTTTACCT
	TCTCCAAACC	AATACATGAA	AACAATACTA	CAGGAAACTT	TTACATTATT
ם המשת המ המ	TACCAAGTAT	TTCACATTTT	GGGAACTTTT	CAGCTAAAGA	AGAAAAAAAC
		2111	Z.Z.V		
スペススペで及べて	GAATTTTTGG	ATTAAAAGAA	TCATGGACTG	GTGGTATCAT	300
		חדפ	/AU		
CAACATGCAG	CTTTTAATAT	CCCAAATTAT	TCATTTAAAT	ATGAAAATAA	TCCATTTIA
CCATTTCCAG	GGGTAATTGG	CTATTCAATA	GGTAGTCCAA	GAATAGAATT	TGAAGTATCA
TACCAGACAT	TCGATGTACA	AAATCCAGGA	GATAAGTTTA	ACAATGATGC	ACATAAGTAT 480
	441		401	,	
ייבייברייייים איניי	CCAATGATT	CAGTAAAACA	ATGAAAAGTO	GTAAATTCGT	TTTTCTCAAA
		A 5.11	1 321	, , , , , , , , , , , , , , , , , , , ,	<i>(</i>
namcaagga'	r TAAGTGACA	T ATCACTCAT	TTAAATGTA	r GTTATGATAT	AATAAACAAA 600
		a 571	יסכ ו	U	•
AGAATGCCT'	T TTTCACCTT	A CATATGTGC	A GGCATTGGT	A CTGACTTAAT	ATTCATGTTT 0 660
			n 54	U 00,	•
GACGCTATA	A ACCATAAAG	C TGCTTATCA	a ggaaaatta	G GTTTTAATT	A TCCAATAAGC 720
		E 0	ก / บ	U / -	•
CCAGAAGCT	A ACATTTCTA	T GGGTGTGCA	C TTTCACAAA	G TAACAAACA	A CGAGTTTAGA 0 780
73					
GTTCCTGTT	-	C TGGAGGACT	C GCTCCAGAT	A ATCTATTIG	C AATAGTAAAG
		na 81	.ი 82	<u> </u>	0 0 0
TTGAGTAT	AT GTCATTTT	GG GTTAGAATT	TT GGGTACAG	G TCAGTTTT	A A
110			Fig. 24A		
			rig. 24A		
			••	40	60
:	10	20		<b>3</b> V -	,0
MNCKKFLI'	TT TLVSLTIL	LP GISFSKPI	HE NNTTGNEY	00 · 1	IF GNFSAKEEKN
•	70.	80	90 1		
TTTGIFGL	KE SWTGGIIL	DK EHAAFNIP	NY SEKYENNE	EL GENGVIGI	SI GSPRIEFEVS 70 180
4	70 1	40 1	50 I	. שם	
		KY CALSNOSS	KT MKSGKEVE 10 2	יסור המה או אור	LM LNVCYDIINK 30 240
1	.90 2	200 2	TO CALCEMAN		•••
		MF DAINHKAA	TO GETTERNIE	280 2	VH FHKVTNNEFR 90 300
∴ 2	250 2	260 2	70 4	.50 2	••
VPVLLTAG	GL APDNLFAI	TVK LSICHFGL	Er GIKVSF.	• • • • • • • • • • • • • • • • • • • •	

Fig. 24B

10	20	30	40	50	60
ATGAATAATA	AACTCAAATT	TACTATAATA	AACACAGTAT	TAGTATGCTT	ATTGTCATTA
70	80	90	100	110	120
CCTAATATAT	CTTCCTCAAA	GGCCATAAAC	AATAACGCTA	AAAAGTACTA	CGGATTATAT
130	140	150	160	170	180
	AATATAAACC	CAGTGTTTCT	GTTTTCAGTA	ATTTTTCAGT	TAAAGAAACC
190	200	210	220	230	240
AATGTCATAA	CTAAAAACCT	TATAGCTTTA	AAAAAAGATG	TTGACTCTAT	TGAAACCAAG
250	260	270	280	290	300
ACTGATGCCA	GTGTAGGTAT	TAGTAACCCA	TCAAATTTTA	CTATCCCCTA	TACAGCTGTA
310	320	330	340	350	360
TTTCAAGATA	ATTCTGTCAA	TTTCAATGGA	ACTATTGGTT	ACACCTTTGC	TGAAGGTACA
370	380	390	400	410	420
AGAGTTGAAA	TAGAAGGTTC	TTATGAGGAA	TTTGATGTTA	AAAACCCTGG	AGGCTATACA
430	440	450	460	47.0	480
CTAAGTGATG	CCTATCGCTA	TTTTGCATTA	GCACGTGAAA	TGAAAGGTAA	TAGTTTTACA
490	500	510	520	530	540
CCTAAAGAAA	AAGTTTCTAA	TAGTATTTT	CACACTGTAA	TGAGAAATGA	TGGATTATCT
550	560	570	580	590	600
ATAATATCTG	TTATAGTAAA	TGTTTGCTAC	GATTTCTCTT	TGAACAATTT	GTCAATATCG
610	620	630	640	650	660
CCTTACATAT	GTGGAGGAGC	AGGGGTAGAT	GCTATAGAAT	TCTTCGATGT	ATTACACATT
670	680	690	700	710	720
AAGTTTGCAT	ATCAAAGCAA	GCTAGGTATI	GCTTATTCTC	TACCATCTAA	CATTAGTCTC
730					
TTTGCTAGTT	TATATTACCA	TAAAGTAATO	GGCAATCAAT	TTAAAAATTI	AAATGTCCAA
790	800	810	820	830	840
CATGTTGCTG	AACTTGCAAG	TATACCTAA	A ATTACATCC	CAGTTGCTAC	ACTTAATATT
850	860	870	880	890	900
GGTTATTTT	GAGGTGAAAT	TGGTGCAAG	A TTGACATTT	AA	• • • • • • • • •
		•			

# Fig. 25A

. 10	20	30	40	50	60
MNNKLKFTII	NTVLVCLLSL	PNISSSKAIN	NNAKKYYGLY	ISGQYKPSVS	VFSNFSVKET
70	80	90	100	110	120
NVITKNLIAL	KKDVDSIETK	TDASVGISNP	SNFTIPYTAV	FQDNSVNFNG	TIGYTFAEGT
130	140		160	170	180
	FDVKNPGGYT	LSDAYRYFAL	AREMKGNSFT	PKEKVSNSIF	HTVMRNDGLS
190	200	210	220	230	240
		PYICGGAGVD	AIEFFDVLHI	K. AYQSKLGI	AYSLPSNISL
250	260	270	280	290	300
FASLYYHKVM			ITSAVATLNI	GYFGGEIGAR	LTF

Fig. 25B

10	20	30	40	50	60
ATGGCAAATT	<del></del> -		CTAATGACAG		ATTATTTCAC
70	80	90	100	110	120
ATGTTATTTC	TACCTCATGT		AAAAATACAA		
130	140	150	160	170	180
TACATCAGTG	GACAGTATAA	CCCTAGTGTT	TCTGTTTTTA	GCAATTTTTC	AGCAAAAGAA
190	200	210	220	230	240
ACCAATGTTC	ATACAGTACA	ACTCATGGCG	CTTAAAAAAG	ACATTGATTC	TATTGAAGTT
250	260	270	280	290	300
GATACTGGAA	ATAGCGCAGG	TATTAGCAAA	CCACAAAATT	TCACAGTTCT	TTATACTCCA
310	320	330	340	. 350	360
AAATTTCAAG	ATAATGTTGC	TGGTCTTAGC	GGTGCACTTG	GATTCTTTTA	TTCTAAAGGA
370	380	390	400	410	420
TTAAGGATTG	AAATGGGGTT	TTCTTATGAA	AAATTTGATG	CTAAAGACCT	TGGTGAGTAC
430	440	450	460	470	480
	AAGATGCTTA	TAGATATTTT	GCTCTAGTAC	GTGAAATGCA	TGTTAGTCTC
490	500	510	520	530	540
			CATTATACTG	TTATGAGAAA	TGATGGTATA
550	560	570	<b>580</b>	- 590	600
TCTATTTCTT		AAATGGCTGC		TTTTCCAGTT	TATCTTTGTC
610	620	630	640	650	660
ACCTATATGT			GCTATAGAAT		
670	680	690	700	710	720
			ACTTATTCTG		
730	740	750	760	770	780
TTTGCAGATG			GGCAATAAAT		
790	800	810	820	830	840
TACGTTAATA	860		GTTACATCTG		
		870	088 TTTATATTTT	890	900
GGCIACCICG	GIGGIGAAAI	TGGCATAAGA	TTTATATTT	AA	• • • • • • • • •
•		Fig	. 26A		
			· -		
10	20	30	40	50	60
MYKKYKLMTA	GVVLFHMLFL	PHVSFAKNTN	SNKLGLYISG	QYNPSVSVFS	NFSAKETNVH
70	80	90	100	110	120
TVQLMALKKD	IDSIEVDTGN	SAGISKPONF	TVLYTPKFQD	NVAGLSGALG	FFYSKGLRIE
130	140	150	160	170	180
MGFSYEKFDA	KDLGEYTKIK	DAYRYFALVR	<b>EMHVSLIYPK</b>	DNNTGTHYTV	MRNDGISISS
190		210	220		240
			LNAYILSLLA	KVVKVLTYSV	
250		270			300
YYHKVMGNKF	KNLPVQYVNT	LEEYPRVTSA	IATLDIGYLG	GEIGIRFIF.	••••••

Fig. 26B

10	20	30	40	50	60
		TAAAAGTCAA	TTCTTAATAA	GATTTATATT	TTTAACATGC
70	80	90	· 100	110	. 120
ATGCTGTCAT	TACCTAATAT	ATCTCTTTCA	AAAGTAAATA	ACGAAAAACA	TTCTGGTTTG
130	140	150	160	170	180
TATATTACCG	GGCAATACAA	ACCCAGTGTT	TCTGTTTTCA	GTAATTTTTC	AGTTAAAGAA
190	200	210	220	230	240
カーにある 一でででで		TCTCATAGCT	CTTAAACAAG	ATGTTGATTC	TGTTGAAATT
250	260	270	280	290	300
CATACTECTA			AACCCATCTA	ACTTTACAAT	CCCTTATACT
310	320	330	340	350	360
		TACTARCTCC	AATGGCTCTA	TTGGTTATGC	TTTTGCTGAA
370	380	390	400	410	420
CCTCCAAGAA			GAAAAATTTG	ATGTTAAAAA	TCCCACAGGG
430	440	450	460	47.0	480
7.57.400.404				CACGTGAAAT	AAATATTTCT
TATACTACAG	500	510		530	540
490				ATGTCGTAAT	GAAAAACGAT
	560		580		600
550		ひてと	ੑ ৾ঌড়ড়ড়৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻৻	ATTTTTCTTT	
GGGTTATCTA					
610				CCATAGAATT	
CCTATATCAC					
670					ATTACGTAAA
	AATTTGCTTA				
730	740	750	,	,	, , , , ,
ATCAACTTAT					TAAAAACCTG
790	800	810			,
					AGTTGCTACA 900
850			-	,	
CTTGATATAC	CATATTTTG(	TAGTGAAGC	r ggcataagai	A TTATATTTI	

# Fig. 27A

60	50	40	UL	ZU	
nfsvætnfh	QYKPSVSVFS	EKHSGLYISG	PNISLSKVNN	FIFLTCMLSL	MNNKSQFLIR
	110	100	90	80	70
GYAFAEGPRI	DNHTNCNGSI	FTIPYTAEFQ	NTAGISNPSN	VDSVEIDTGS	TKHLIALKQD
180	170	160	150	140	130
VVMKNDGLSI	KQKEGSGIYH	REINISLFQP	KDAYRYFALA	VKNPTGYTTV	EIELSYEKFD
240	230	220	210	200	190
YQLLRKINLF	FAYOSKAGIS	IEFFDALHVK	YLCGGMGINA	FSLNNLPISP	LSNIVNICYD
300	290	280	270	260	250
IF	YFGSEAGIRI	TSAVATLDIA	VHELKDNPKV	NKFKNIKVOH	IDVYYYEVIS

Fig. 27B

			40	E0.	60
10	20	30	40	50	~ -
•				TAATATGCTT 110	120
70	80	90	100		
				AACATTCTGG 170	180
130	140	150	160		
AGCGGACAAT				TTTCAGTAAA	240
190	200	210	220	230	TATGAACATC
		AGCTCTTAAA		ATTCTATTTC 290	300
250	260	270	280		• • • • • • • • • • • • • • • • • • • •
				ATCTTCCTTA	360
310	320	. 330	340	350	
				ATTCACTTTT	TGAACAACTA
370	380	390	400	410	
			•	AAAATCCTGG	
430	440	450	460	47.0	480
TTAAATGATG				TGGGACAAGA	
490	500	510	520	530	540
AATAAGCATC				AAACATATTA	
550	560	570	580	590	600
AGAAATAATG	GGTTATCTAT			GCTGCTATAA	
610	620	630	640	650	. 660
AATGATTTAT				GTGTAGATGC	
670	680	690	700	710	720
TTTGATGCAC	TGCATCTTAA			TAGGAGCTAC	
730	740	750	760	770	780
TCAGACAACA	TTAGTTTATT	TACAAATGGA		AAGTAATAGG	
790		810			840
AAAAACTTAA	. AAGTCCAATA	TATAGGTGAA	CTTAAAGAGA	ACCCGAAAAT	
850		870	• • •	890	900
GTTGCTACTC	: TCAATGTTGG	ATACTTTGGA	GGTGAAATTG	GAGTAAGACT	
910	920	930	940	950	960
		• • • • • • • • •	• • • • • • • • •	• • • • • • • • •	• • • • • • • • •
		Fig	. 28A		
		rig	. 20A		
. 10	20	30	40	50	60
				SGQYKPSVSI	FSKFSVKETN
70.		90			120
				FQDNAFNFSG	AIGYSLFEQL
	140	150			
				NKHLSPKEEH	
190		210		230	
				FDALHLKLAL	•
250	•				
				VATLNVGYFG	•
PONTOTEING	TIUMATANAE	""INVANTAGE	TUBRENTON	***************************************	

Fig. 28B

10	20	30	40	50	60
AAGCTTCTTA	TGAAGAATTT	GACGTTAAAA	ATCCTGAAGG	ATCTACTACA	GACTCCTATA
70	80	. 90	100	110	120
GATATTTCGC	GTTAGCACGT	GGCATGGATG	GTAATAATAT	TCCTACAAGT	CAAAAATTTA
130	140	150	160	170	180
CTGTAATGAG	AAACGACGGG	TTATTAATCT	CATCTGTTAT	GATAAATGGC	TGTTACAATG
190	200	210	220	230	240
TCATACTAAA	TGATATACAA	GCAGAACCTT	ACATATGTGC	AGGACTAGGA	GGAGATTTTA
250	260	270	280	290	300
TAGAATTCTT	CAATGGCTTT	CATGTTAAGC	TAGCTTATCA	AGGTAAAGTA	GGCATTAGTT
310	320	330	340	350	360
ATCAAATATT	CCCTGAAGTA	AGATTATTTA	TTGATGGATA	CTACCATAAA	GTAAAAGGCA
370	380	390	400	410	420
ACAAGTTTAA	AAATTTACAC	GTTCAACATG	TAGGTGCACT	TGCAGCACTC	CCTAAAGTTA
430	440	450	460	470	480
CATCTGCAGT	TGCAACACTT	AATATTGGAT	ACTTTGGTTG	TGAAGCTGGA	GTAAGATTCA
490	500	510	520	530	540
TATTTTAA	• • • • • • • • •	• • • • • • • • •	• • • • • • • • •	••••••	

# Fig. 29A

60	50	40	30	20	10
SVMINGCYNV	VMRNDGLLIS	NNIPTSQKFT	YFALARGMDG	PEGSTTDSYR	ASYEEFDVKN
120	110	100	. 90	80	
DGYYHKVKGN	QIFPEVRLFI	AYQGKVGISY	EFFNGFHVKL	ICAGLGGDFI	ILNDIQAEPY
180	. 170	1.60	150	140	130
	F	FGCEAGVRFI	SAVATLNIGY	GALAALPKVT	KFKNLHVOHV

Fig. 29B

10	20	30	40	50	60
ATGAATTATA	AGAAAATTCT	AGTAAGAAGC	GCGTTAATCT	CATTAATGTC	AATCTTACCA
70	80	90	100	110	. 120
TATCAGTCTT	TTGCAGATCC	TGTAGGTTCA	AGAACTAATG	ATAACAAAGA	AGGCTTCTAC
130	140	150	160	170	180
ATTAGTGCAA	AGTACAATCC	AAGTATATCA	CACTTTAGAA	AATTCTCTGC	TGAAGAAACT
190	200	210	220	230	240
CCTATTAATG	GAACAAATTC	TCTCACTAAA	AAAGTTTTCG	GACTAAAGAA	AGATGGTGAT
250	260	270	280	290	
ATAACAAAAA	AAGACGATTT		GCTCCAGGCA	TTGATTTTCA	AAATAACTTA
310	320	330	. 340	350	360
ATATCAGGAT	TTTCAGGAAG	TATTGGTTAC	TCTATGGACG	GACCAAGAAT	AGAACTTGAA
	380	390	400	410	420
370	• • •				TGGTGAATAC
GCTGCATATC	ACAATTTAAT	CCAAAAACAC	_		
430	440	450	460	470	480
TATAAACATT	·TTGCATATCT	CGTAAAGATG	CCATGGAAGA	TCAGCCATAT	GTTGTTCTTA
490	500	510	520	530	540
AAAATGACGG	CATAC		• • • • • • • • •		• • • • • • • • •

# Fig. 30A

	20	30	40	50	60
10	20	woods Device	PUNDNEEGEY	ISAKYNPSIS	HFRKFSAEET 120
MNYKKILVRS	ALISLMSILP	YQSFADPVGS	100	110	120
70	. 80	90			SMOGPRIELE
PINCTUSLIK	KVFGLKKDGD	ITKKDDFTRV	APGIDFQNNL	ISGESGSIGI	SMDGPRIELE 180
130	140	150	100	170	
HAUT LUNGS	DNNDTDNGEY	YKHFAYLVKM	PWKISHMLFL	KMTAY	• • • • • • • • •

Fig. 30B

		HV1
OMP-1F CMF-1B CMF-1D CMF-1C CMF-18 F28 MAP-1 CMF-1A	SV	
CMP-1F CMP-1E CMP-1D CMP-1C CMP-19 P28 HAP-1 CMP-1A	YSTATEONPP LGFAGAVGYL MOMPRIELEM SYNTTDVRNQ CHRYDDDAN - RYVALTH- MSGCHEMAG DEFVFLONES  LS G.V.F.V - R.C. OG - QURBGIERT S.Y.L. S.  LS G.F.V A G - R.C. OR - RASSTRANT SHY.L.  PALEFQ. LI S.S.I. A D. A AVCK A. P D. DT. SCHY Y. FG. SR - RADALS S. MS. L.  S P.V R.F. G - R.C. SH - AADMS.S MS.  HV3	117MV.TITAV. 188 
CHP-17 CHP-18 CHP-10 CHP-16 CHP-18 P28 HAP-1	FSFYICADVG TOLISGEAI MPRISYQGEL GLSYSISPEA SVFVGGGFRK VIGNEFEDIP AMIPSTSTLT GE-HF	A T

Fig. 31

International application No. PCT/US98/19600

	SSIFICATION OF SUBJECT MATTER		
	: A01N 43/04; A61K 39/02 : 514/44; 424/234.1		
	to International Patent Classification (IPC) or to both	national classification and IPC	
B. FIEL	DS SEARCHED		
Minimum d	ocumentation searched (classification system followed	d by classification symbols)	
<b>U.S.</b> :	514/44; 424/234.1		
Documental	tion searched other than minimum documentation to the	extent that such documents are included	in the fields searched
Pleatennia	data base consulted during the international search (na	ame of data have and where practicable	search terms used)
APS, DI		une of data base and, where practicable	, search willis uses,
	ms: erlichi?, protein?, antigen?, polypeptide?, dna, r	ecombinant?, clone?, dna, polynucleoti	de, nucleotide?
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ap	propriate, of the relevant passages	Relevant to claim No.
A	US 5,789,176 A (DAWSON et al) 04 August 1998, see abstract, claims and entire document.		1, 9, 11, 19, 21- 22
A	US 5,401,656 A (DAWSON et al) 28 March 1995, see abstract, claims and entire document.		1, 9, 11, 19, 21- 22
A	US 5,413,931 A (DAWSON et al) claims and entire document.	09 May 1995, see abstract,	1, 9, 11, 19, 21- 22
Y,E	US 5,869,335 A (MUNDERLOH et abstract, claims and entire document.	al) 09 February 1999, see	1, 9
		•	
X Furt	her documents are listed in the continuation of Box C	See patent family annex.	
<ul> <li>Special categories of cited documents:</li> <li>'I' later document published after the international filing date or priority date and not in conflict with the application but cited to understand</li> </ul>			
	cument defining the general state of the art which is not considered be of particular relevance	the principle or theory underlying th	
	Compared to the commerce of th		
ci	*L° document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other process (see section).  *Y° document of particular relevance; the claimed invention cannot be		
*O* do	considered to involve an inventive step when the document is		
*P* document published prior to the international filing date but later than *& document member of the same patent family the priority date claimed			
	e actual completion of the international search	Date of mailing of the international se	arch report
18 FEBR	UARY 1999	40 LTD 1999	
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	n, D.C. 20231	GINNY PORTNER	1
l Bandanita N	Va (703) 305 3330	Telephone No. (703) 308-0196	v ~17

International application No. PCT/US98/19600

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 2-8, 10, 12-18, 20 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
The claims as submitted evidenced blank lines, therefore the claims were incomplete and found to be unsearchable.
Claims Nos.:  because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
·
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

International application No. PCT/US98/19600

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X  Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). BROUQUI, P. et al. 'Serologic diagnosis of human monocytic ehrlichiosis by immunoblot analysis'. Clinical Diagnostic Laboratory Immunology, November 1994, Vol. 1, No. 6, pages 645-649, see entire abstract.	11,19, 21, 22  1, 9
<b>Y</b> .	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). BROUQUI, P. et al. 'Antigenic characterization of ehrlichiae: protein immunoblotting of Ehrlichia canis, Ehrlichia sennetsu, and Ehrlichia risticii'. Journal of Clinical Microbiology. May 1992, Vol. 30, No. 5, pages 1062-1066, see entire abstract.	19, 21, 22
X  Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). CHEN, SM et al. Identification of the antigenic constituents of Ehrlichia chaffeensis'. American Journal of Tropical Medicine and Hygiene. January 1994, Vol. 50, No. 1, page 52-58, see entire abstract.	11, 21  1
х  Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). CHEN, SHENG-MIN et al. 'Analysis and Ultrastructural localization of Ehrlichia chaffeensis proteins with monoclonal antibodies'. The American Journal of Tropical Medicine and hygiene. April 1996, Vol. 54, No. 4, pages 405-412, see entire abstract.	11, 19 21, 22  1
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<b>Y</b>	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). DAWSON, JE et al. 'The Interface between research and the diagnoses of an emerging tick-borne disease, human ehrlichiosis due to Ehrilichia chaffeensis'. Archives of Internal Medicine, 22 January 1996, Vol. 156, No. 2, pages 137-end, see entire document.	1, 9
Y	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). KELLY, PJ et al. 'Serological evidence for antigenic relationships between Ehrlichia canis and Cowdria ruminantiu'. Research in Veterinary Science. March 1994, Vol. 56, No. 2, page 170 174, see entire abstract.	19

International application No. PCT/US98/19500

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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?	Database Medline on Dialog, US National Library of Medicine (Bethesda, MD, USA,). YU, XJ et al. 'Sequence and characterization of an Ehrlichia chaffeensis gene encoding 314 amino acids highly homologous to the NAD A enzyme'. FEMS Microbiology Letters, 01 September 1997, Vol. 154, No. 1, pages 53-58, see entire document.	1, 9
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